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(54) SUBLINGUAL FENTANYL SPRAY

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- USPC 514/329; 128/200.14; 424/400; 514/506;
- (58) Field of Classification Search See application file for complete search history.

(56)**References Cited**

U.S. PATENT DOCUMENTS

4,244,478	\mathbf{A}	1/1981	Handman
5,219,083	\mathbf{A}	6/1993	Liebert et al.
5,958,379	A	9/1999	Regenold et al.
5,976,504	\mathbf{A}	11/1999	Russell
6,759,059	В1	7/2004	Pettersson et al.
2002/0055496	$\mathbf{A}1$	5/2002	McCoy et al.
2002/0160991	$\mathbf{A}1$	10/2002	Shao
2003/0039680	$\mathbf{A}1$	2/2003	Dugger, III
2003/0077228	$\mathbf{A}1$	4/2003	Dugger, III
2003/0077229	$\mathbf{A}1$	4/2003	Dugger, III
2003/0082107	$\mathbf{A}1$	5/2003	Dugger, III
2003/0095925	$\mathbf{A}1$	5/2003	Dugger, III
2003/0095926	$\mathbf{A}1$	5/2003	Dugger, III
2003/0095927	$\mathbf{A}1$	5/2003	Dugger, III
2003/0190286	$\mathbf{A}1$	10/2003	Dugger, III
2003/0190290	A1*	10/2003	Ross 424/45
2004/0092428	$\mathbf{A}1$	5/2004	Chen et al.
2004/0120895	$\mathbf{A}1$	6/2004	Dugger, III
2004/0136913	$\mathbf{A}1$	7/2004	Dugger, III et al.
2004/0136914	$\mathbf{A}1$	7/2004	Dugger, III et al.
2004/0136915	$\mathbf{A}1$	7/2004	Dugger, III et al.

2004/0141923	A1	7/2004	Dugger, III et al.
2004/0265239	A1	12/2004	Dugger, III et al.
2005/0002867	A1	1/2005	Dugger, III et al.
2005/0163719	A1	7/2005	Dugger, III et al.
2005/0180923	A1	8/2005	Dugger, III et al.
2005/0281752	A1	12/2005	Dugger, III
2005/0281753	A1	12/2005	Dugger, III
2005/0287075	A1	12/2005	Dugger, III
2006/0062812	A1*	3/2006	Ross et al 424/400
2009/0124554	A1	5/2009	Dugger, III
2009/0162300	A1	6/2009	Dugger, III et al.
2009/0176834	A1*	7/2009	Kottayil et al 514/329

FOREIGN PATENT DOCUMENTS

RU	2156126	9/2000
RU	2232580	7/2004
WO	WO 90/07333	7/1990
WO	WO 00/47203	8/2000
WO	WO 01/97780 A2	12/2001
WO	WO 2004/016243 A2	2/2004
WO	WO 2004/080382	9/2004
WO	WO 2004/075877	10/2004
WO	WO 2007-007059	1/2007

OTHER PUBLICATIONS

Smyth, et al, 2003, AAPS PharmSciTech 2003; 4 (3): 1-11.* Smyth & Hickey (2003) Multimodal Particle Size Distributions Emitted From HFA-134a Solution Pressurized Metered-Dose Inhal-

ers. AAPS PharmSciTech 2003; 4 (3) Article 38. pp. 1-11.* International Search Report and Written Opinion dated Jul. 11, 2008 from corresponding Int'l Application No. PCT/US07/02163.

Mathar, L.E., et al. "Pulmonary administration of aerosolised fentanyl: pharmacokinetic analysis of systemic delivery" Br. J. Clin. Pharmacol. Jan. 1998, vol. 46, pp. 37-43.

Marier, J-F., et al. "Comparative bioequivalence study between a novel matrix transdermal delivery system of fentanyl and a commercially available reservoir formulation", Br. J. Clin. Pharmacol. Aug. 2006, vol. 63, No. 1, pp. 121-124, esp. p. 123, Figure 1.

International Preliminary Report on Patentability dated Sep. 4, 2008 from corresponding Int'l Application No. PCT/US07/02163.

Examination Report dated Aug. 6, 2009 from corresponding Australian Application No. 2007208229

Examination Report dated May 5, 2010 from corresponding Canadian Application No. 2,637,672.

Examination Report dated Mar. 18, 2010 from corresponding New Zealand Application No. 569949.

An English translation of the Russian Examination Report dated Aug. 2009 from corresponding Russian Application No. 2008130763. ISR and Written Opinion from related Int'l Application No. PCT/

US08/09359 dated Jan. 9, 2009. Int'l Preliminary Report on Patentability from related Int'l Applica-

tion No. PCT/US08/09359 dated Feb. 2, 2010.

(Continued)

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(57)ABSTRACT

The present invention is directed to sublingual formulations containing fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, suitable for administration to a patient, and methods for treatment with the formulations.

3 Claims, 7 Drawing Sheets

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OTHER PUBLICATIONS

Lejus, et al., "Fentanyl versus sufentanil: plasma concentrations during continuous epidural postoperative infusion in children," British Journal of Anaesthesia, vol. 85, Issue 4, Oct. 2000, pp. 615-617. Office Action on Patent Application No. 200780003555.X dated Jul. 9, 2010.

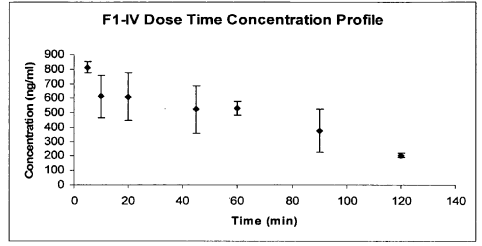
Examination Report for corresponding European Patent Application No. 07762549.9, mailed on Nov. 30, 2010.

Supplementary Search Report for corresponding European Patent Application No. 97762549.9, mailed on Nov. 30, 2010.

* cited by examiner

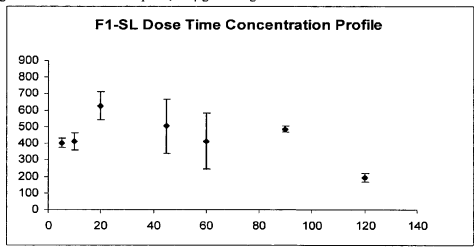
Plasma concentration-time curves after IV & SL doses of Fentanyl

Figure 1: Formula of Example 1, 50 µg IV dose



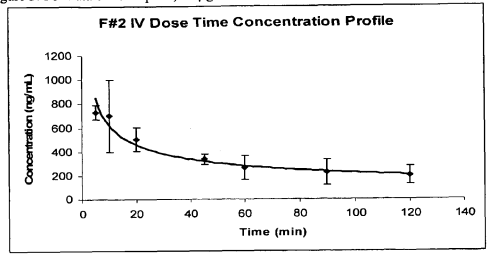
Mean (\pm S.E.) plasma concentration—time profiles following intravenous administration of Fentanyl (n=3).

Figure 2: Formula of Example 1, 50 µg Sublingual dose



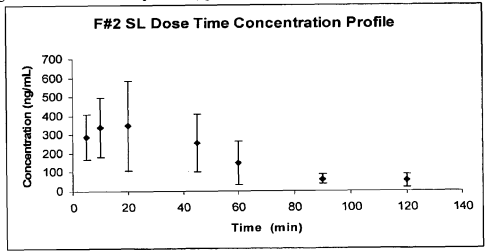
Mean (\pm S.E.) plasma concentration–time profiles following sublingual administration of Fentanyl (n=3)

Figure 3: Formula of Example 2, 80 µg IV dose



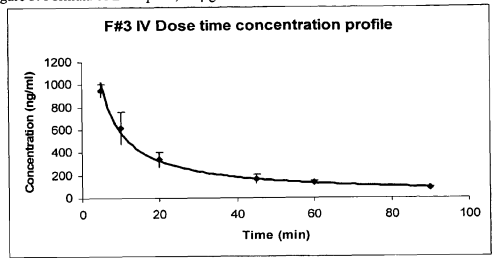
Mean (\pm S.E.) plasma concentration-time profiles following intravenous administration of Fentanyl (n=3).

Figure 4: Formula of Example 2, 80 µg SL dose



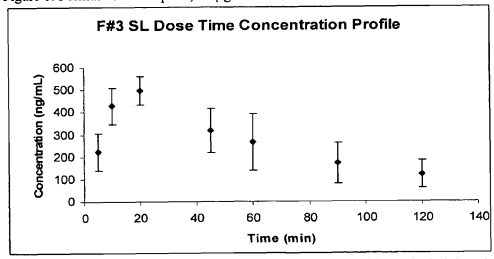
Mean (\pm S.E.) plasma concentration—time profiles following sublingual administration of Fentanyl (n=3)

Figure 5: Formula of Example 3, 50 µg IV dose



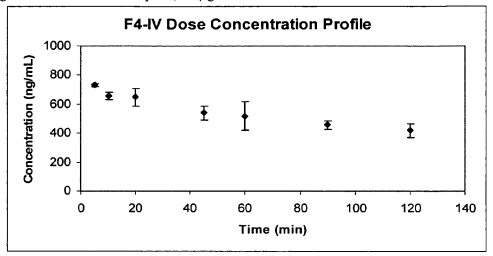
Mean (\pm S.E.) plasma concentration—time profiles following intravenous administration of Fentanyl (n=3)

Figure 6: Formula of Example 3, 50 μg SL dose



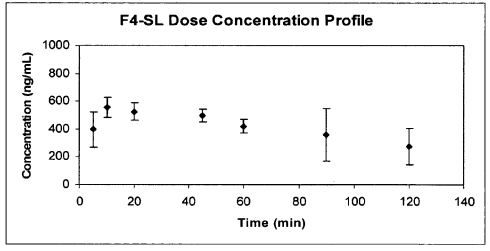
Mean (\pm S.E.) plasma concentration—time profiles following sublingual administration of Fentanyl (n=3)

Figure 7: Formula of Example 4, 50 µg IV dose



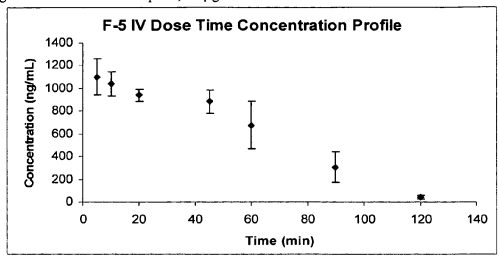
Mean (\pm S.E.) plasma concentration—time profiles following intravenous administration of Fentanyl (n=3)

Figure 8: Formula of Example 4, 50 µg SL dose



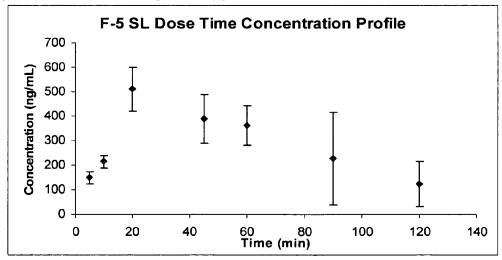
Mean (\pm S.E.) plasma concentration—time profiles following sublingual administration of Fentanyl (n=3)

Figure 9: Formula of Example 5, 50 µg IV dose



Mean (\pm S.E.) plasma concentration–time profiles following intravenous administration of Fentanyl (n=3)

Figure 10: Formula of Example 5, 50 µg SL dose



Mean (\pm S.E.) plasma concentration—time profiles following sublingual administration of Fentanyl (n=3)

Figure 11: Vignetting Results Dv10, Dv50, Dv90 vs. Distance from Range Lens

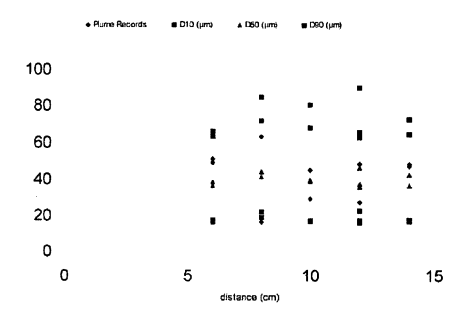
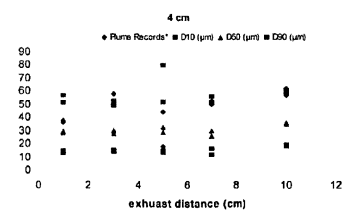
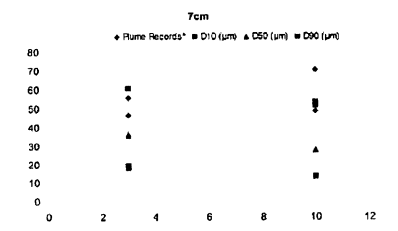


Figure 12: Exhaust Results Dv10, Dv50, Dv90 vs. Distance to Exhaust at 4 cm Device to Laser Beam



Note: 10 cm distance used for plotting of results with no exhaust fan present. Reference: Notebook 1, LDR 108

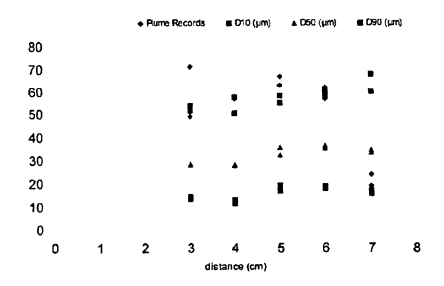
Figure 13: Exhaust Results Dv10, Dv50, Dv90 vs. Distance to Exhaust at 7 cm Device to Laser Beam



Note: 10 cm distance used for plotting of results with no exhaust fan present.

Reference: Notebook 1, LDR 108

Figure 14: Device to Laser Beam Placement Results, Dv10, Dv50, Dv90 vs. Distance to Laser Beam



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SUBLINGUAL FENTANYL SPRAY

FIELD OF THE INVENTION

The invention is directed to sublingual formulations containing fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, suitable for administration to humans, and methods for treatment with the sublingual formulations.

BACKGROUND OF THE INVENTION

Fentanyl is a μ -opioid receptor agonist with analgesic potency approximately 80-100 times that of morphine. In clinical settings, fentanyl exerts its principal pharmacologic effects on the central nervous system. Its primary actions are 15 analgesic and sedation.

The analgesic effects of fentanyl are related to the blood level of the drug. In general, the minimum effective concentration and the concentration at which toxicity occurs rise with increasing tolerance to any and all opioids. The rate of 20 development of tolerance may vary widely among individuals. All opioid mu-receptor agonists, including fentanyl, produce dose dependent respiratory depression. The risk of respiratory depression is typically less in patients receiving chronic opioid therapy who develop tolerance to respiratory depression and other opioid effects. Serious or fatal respiratory depression can occur, even at recommended doses, in vulnerable individuals.

Orally administered fentanyl is subject to first pass effect metabolism as upwards of 50% or more of orally adminis- 30 tered fentanyl is not absorbed. Other forms of delivery such a parenteral, buccal, and transdermal have been utilized to decrease or avoid this first pass effect for fentanyl.

Fentanyl is currently available in injectable form, as a lozenge (e.g. Actiq®), and as a transdermal system (e.g., 35 Duragesic® 25, 50, 75, and 100 µg of fentanyl per hour). Duragesic® provides continuous systemic delivery of fentanyl for approximately 72 hours. Duragesic® is indicated in the management of chronic pain in patients requiring continuous opioid analgesia for pain that is not optimally managed with lesser means such as acetaminophen-opioid combinations, non-steroidal analgesics, or prn (as needed) dosing with short-acting opioids. Duragesic® is typically not suitable for patients experiencing acute pain due to the delay in absorption of the fentanyl through the patch, or postoperative pain 45 because serious or life-threatening hypoventilation could result.

Actiq® is a solid formulation of fentanyl citrate, intended for oral transmucosal administration. Actiq® is a lozenge attached to a handle similar in shape to a lollipop. The handle 50 is purportedly to allow the Actiq® unit to be removed from the mouth if signs of excessive opioid effects appear during administration. Actiq® is indicated for the management for breakthrough cancer pain in patients with malignancies who are already receiving and who are tolerant to opioid therapy 55 for their underlying persistent cancer pain. Actiq® is contraindicated in the management of acute or postoperative pain.

Sublingual tablets and lozenges (e.g., Actiq®) which may be used for acute pain or breakthrough pain have certain 60 disadvantages. A disadvantage, amongst others, is that after intake the active agent in these pharmaceutical agents must first be released and dispersed prior to being available for resorption in dissolved form. In addition, the absorption pharmacokinetics of fentanyl from Actiq® may vary depending 65 on the fraction of the dose that is absorbed through the oral mucosa and the fraction swallowed. Further, certain lozenges

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may be in the form of a candy which require medical supervision and may be socially questionable.

There exists a need in art for a sublingual formulation including fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, which is suitable for effective pain management.

SUMMARY AND OBJECTS OF THE INVENTION

It is an object of the invention to provide a fentanyl formulation suitable for sublingual administration for effective pain management.

It is an object of certain embodiments of the invention to provide methods and compositions capable of rapidly inducing a state of sedation, analgesia, and/or anesthesia.

It is a further object of certain embodiments of the invention to provide methods and compositions for fentanyl administration which minimize the underdosing and/or overdosing of a patient in need of fentanyl therapy.

It is a further object of certain embodiments of the invention to provide methods and compositions suitable for the treatment of breakthrough pain in patients receiving chronic pain treatment.

It is a further object of certain embodiments of the present invention to provide a method for sublingual administration of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, in a controlled amount for the treatment of pain.

It is a further object of certain embodiments of the present invention to provide a dosage form of an opioid analgesic which can be administered sublingually in a manner which will cause substantial sublingual absorption without substantial risk of the dose passing into the lungs of the recipient.

The above-mentioned objects and others are achieved by virtue of the present invention, which is directed in part to a method for sublingually administering fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, to provide fast-acting relief in a formulation in which a substantial portion of the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof will not be passed into the lungs of the patient.

In certain embodiments the present invention is directed to a sublingual fentanyl formulation comprising discrete liquid droplets comprising an effective amount of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, said droplets having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more preferably a mean diameter of from about 20 to about 200 microns.

In certain embodiments, the present invention is directed to a sublingual fentanyl formulation comprising discrete liquid droplets of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof; in a pharmaceutically acceptable liquid carrier; said droplets having a size distribution of from about 5 microns to about 500 microns, preferably from about 10 microns to about 200 microns, preferably from about 20 microns to about 100 microns, more preferably from about 30 microns to about 70 microns.

In certain preferred embodiments, none of the particles have a diameter which would allow the fentanyl, pharmaceutically acceptable salt thereof, or derivative thereof to be delivered to the lung upon sublingual administration.

In certain embodiments, the present invention is directed to a unit dose of a sublingual fentanyl formulation, said unit dose comprising discrete liquid droplets of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof; and a pharmaceutically acceptable liquid carrier; said drop-

lets having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more preferably a mean diameter of from about 20 to about 200 microns.

In certain embodiments, the present invention is directed to a unit dose of a sublingual fentanyl formulation, said unit 5 dose comprising discrete liquid droplets of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof; and a pharmaceutically acceptable liquid carrier; said droplets having a size distribution of from about 5 microns to about 500 microns, preferably from about 10 microns to about 200 microns, preferably from about 20 microns to about 100 microns, more preferably from about 30 microns to about 70 microns.

In certain embodiments, the present invention is directed to a method of treating pain comprising sublingually adminis- 15 tering a liquid spray formulation in the form of discrete liquid droplets having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more preferably a mean diameter of from about 20 to about 200 microns, to a human patient experiencing pain, said liquid spray formulation com- 20 prising an effective amount of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, dispersed in a pharmaceutically acceptable liquid carrier.

In certain embodiments, the present invention is directed to a method of treating pain comprising sublingually adminis- 25 a method of treating breakthrough pain comprising sublintering a liquid spray formulation in the form of discrete liquid droplets having a size distribution of from about 5 microns to about 500 microns, preferably from about 10 microns to about 200 microns, preferably from about 20 microns to about 100 microns, more preferably from about 30 microns to 30 about 70 microns to a human patient experiencing pain; said liquid spray formulation comprising an effective amount of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, dispersed in a pharmaceutically acceptable liquid carrier.

In certain embodiments, the present invention is directed to a device which includes a reservoir containing a unit dose of a liquid formulation comprising an effective amount of fentanyl, a pharmaceutically acceptable salt thereof, or derivathe device having an actuator which when actuated delivers the unit dose of the liquid formulation in the form of liquid droplets having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more preferably a mean diameter of from about 20 to about 200 microns. Preferably, 45 the device delivers a therapeutically effective dose of the liquid formulation in the form of liquid droplets having a size distribution of from about 5 microns to about 500 microns, preferably from about 10 microns to about 200 microns, preferably from about 20 microns to about 100 microns, more 50 preferably from about 30 microns to about 70 microns.

In certain embodiments, the present invention is directed to a multi-dose device which includes a reservoir containing a liquid formulation comprising fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof in a pharmaceu- 55 tically acceptable liquid carrier; the device having an actuator which when actuated delivers a therapeutically effective dose of the liquid formulation in the form of liquid droplets having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more preferably a mean diameter of 60 from about 20 to about 200 microns. Preferably, the device delivers a therapeutically effective dose of the liquid formulation in the form of liquid droplets having a size distribution of from about 5 microns to about 500 microns, preferably from about 10 microns to about 200 microns, preferably from 65 about 20 microns to about 100 microns, more preferably from about 30 microns to about 70 microns.

In certain embodiments, the present invention is directed to a method of treating pain comprising utilizing a spray device which includes a reservoir including a liquid formulation comprising fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof in a pharmaceutically acceptable liquid carrier; and an actuator which upon actuation delivers a therapeutically effective amount of liquid droplets to be sprayed from the device having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more preferably a mean diameter of from about 20 to about 200 microns.

In certain embodiments, the present invention is directed to a method of treating pain comprising utilizing a spray device which includes a reservoir including a liquid formulation comprising fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof; and a pharmaceutically acceptable liquid carrier; and an actuator which upon actuation delivers a therapeutically effective amount of liquid droplets having a size distribution of from about 5 microns to about 500 microns, preferably from about 10 microns to about 200 microns, preferably from about 20 microns to about 100 microns, more preferably from about 30 microns to about 70 microns.

In certain embodiments, the present invention is directed to gually administering a liquid spray formulation comprising an effective amount of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, dispersed in a pharmaceutically acceptable liquid carrier to a human patient to treat breakthrough pain experienced by said human patient.

In certain embodiments, the present invention is directed to a method of treating breakthrough pain comprising sublingually administering a liquid spray formulation comprising an effective amount of fentanyl, a pharmaceutically accept-35 able salt thereof, or derivative thereof, dispersed in a pharmaceutically acceptable liquid carrier to a human patient who is receiving chronic pain treatment, and is experiencing breakthrough pain.

In certain embodiments, the present invention is directed to tive thereof in a pharmaceutically acceptable liquid carrier; 40 a method of reducing patient to patient variability for the treatment of breakthrough pain, comprising sublingually administering to a human patient experiencing breakthrough pain a dose of fentanyl in a liquid spray formulation comprising fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, and a pharmaceutically acceptable liquid carrier said liquid spray formulation being administered as discrete liquid droplets having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more preferably a mean diameter of from about 20 to about 200 microns.

> In certain embodiments, the present invention is directed to a method of reducing patient to patient variability for the treatment of breakthrough pain, comprising sublingually administering to a human patient experiencing breakthrough pain a dose of fentanyl in a liquid spray formulation comprising fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, and a pharmaceutically acceptable liquid carrier said liquid spray formulation being administered as discrete liquid droplets having a size distribution of from about 5 microns to about 500 microns, preferably from about 10 microns to about 200 microns.

> In certain preferred embodiments, the liquid spray formulation further includes a pharmaceutically acceptable solvent. Preferably the pharmaceutically acceptable solvent is an organic solvent which is included in an amount suitable for dissolving the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof.

In certain preferred embodiments the formulations of the present invention provide a mean time to maximum plasma concentration (T_{max}) of fentanyl at from about 5 minutes to about 120 minutes, after sublingual administration to humans.

In certain preferred embodiments the formulations of the present invention provide a mean maximum plasma concentration (C_{max}) of fentanyl of about 127 pg/ml to about 213 pg/ml per 100 µg fentanyl after sublingual administration to humans.

In certain preferred embodiments of the present invention the formulations of the present invention do not include a propellant.

In certain embodiments, the formulations of the present invention are suitable for transmucosal administration, 15 including, for example, buccal administration.

In certain embodiments, the present invention is further directed to a method of transmucosally administering fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, to a human to provide fast-acting relief in a formulation in which a substantial portion of the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof will not be passed into the lungs of the patient. In certain preferred embodiments, the transmucosal area is the buccal area of a human

In certain embodiments, the present invention is further directed to the use of a formulation as defined above for the manufacture of a medicament for use as an analgesic, for the treatment of acute pain and/or breakthrough pain, as an anesthetic premedication, for the induction of anesthesia, as a 30 sedative and/or for the treatment of anxiety.

The invention is also directed to a sublingual fentanyl formulation comprising discrete liquid droplets of an effective amount of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof; in a pharmaceutically acceptable liquid carrier; said droplets having a mean diameter of at least about 10 microns, and upon administration to a human patient, at least about 90% of the discrete liquid droplets have a mean diameter equal or greater than about 9 μ m. In other embodiments, not more than about 5% of the discrete liquid 40 droplets have a mean diameter less than 9 μ m. In still other embodiments, the formulation provides a respirable dose of not more than about 5% of the total fentanyl dose contained.

The invention is also directed to a method of treating pain comprising sublingually administering a liquid spray formulation in the form of discrete liquid droplets having a mean diameter of at least about 10 microns to a human patient experiencing pain and at least about 90% of the discrete liquid droplets have a mean diameter equal or greater than about 9 µm upon administration to a human patient, said liquid spray formulation comprising an effective amount of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, dispersed in a pharmaceutically acceptable liquid carrier. In certain other embodiments, not more than about 5% of the discrete liquid droplets have a mean diameter less 55 than 9 µm. In other embodiments, the formulation provides a respirable dose of not more than about 5% of the total fentanyl dose contained.

The invention is also directed to a unit dose or bi-dose device for sublingual administration of a drug comprising: a reservoir containing a unit dose or a bi-dose of a liquid formulation comprising an effective amount of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof in a pharmaceutically acceptable liquid carrier; and

the device having an actuator which when actuated delivers 65 the unit dose of the liquid formulation in the form of liquid droplets having a mean diameter of at least about 10 microns,

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and wherein upon administration to a human patient, at least about 90% of the discrete liquid droplets have a mean diameter equal or greater than about 9 µm.

In other embodiments, not more than about 5% of the discrete liquid droplets have a mean diameter less than 9 μ m. In still other embodiments, the formulation provides a respirable dose of not more than about 5% of the total fentanyl dose contained.

Many patients with e.g., cancer, typically continue to experience moderate to severe pain despite chronic analgesic therapy and this can occur as intermittent breakthrough pain, often due to increases in a patient's activity level. Attempts to counteract this type of pain by increasing the dose of longacting formulations of analgesics often produce slow onset of analgesia and unwanted side-effects of sedation, constipation or nausea and vomiting. However, in certain embodiments the present invention is directed to a formulation which preferably provides a rapidly acting, potent analgesic which reduces the pain, such as breakthrough pain, for the required time and then preferably wears off fairly quickly thereby minimizing the side-effects of the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof.

For purposes of the present invention, derivatives of fentanyl include sufentanil, carfentanil, lofentanil, alfentanil, or the like.

For purposes of the present invention, "breakthrough pain" refers to a pain that exceeds a threshold in a patient which causes cognizable discomfort wherein the pain experienced by the patient is otherwise typically controlled e.g., by chronic analgesic therapy, and tolerated. For example, pain related to medical illnesses, such as cancer, typically fluctuates, and patients often report the experience of cognizable discomfort (e.g., breakthrough pain). Typically breakthrough pain is currently treated with immediate release oral dosage forms which may take up to about 45 minutes or longer for the drug to be absorbed and may result in a delay of the relief of breakthrough pain, as opposed to a liquid spray formulation of the present invention which begins to provide relief of the breakthrough pain almost immediately after administration.

For purposes of the present invention, "chronic pain treatment" refers to a daily or round-the-clock pain treatment. Chronic pain treatment can be oral, parenteral, transdermal, or other suitable means of administration.

For purposes of the present invention, "sublingual" is defined herein as beneath or concerning the area under the tongue.

For purposes of the present invention the term "sublingual administration" is defined herein as the therapeutic administration of a pharmaceutical composition under the tongue.

For purposes of the present invention an "effective amount" of a drug is an amount effective to demonstrate a desired activity of the drug. According to the instant invention, a therapeutically effective amount of fentanyl, pharmaceutically acceptable salt thereof, or derivative thereof, is an amount effective to treat, e.g., noticeably reduce, pain in a patient.

For purposes of the present invention the terms droplets and particles may be used interchangeably.

BRIEF DESCRIPTION OF THE DRAWINGS

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FIG. 1 depicts the mean (\pm S.E.) plasma concentration-time profiles following intravenous administration of Fentanyl of Example 1 (n=3) in the study of Example 6.

FIG. 2 depicts the mean (±S.E.) plasma concentration-time profiles following sublingual administration of Fentanyl of Example 1 (n=3) in the study of Example 6.

FIG. 3 depicts the mean (±S.E.) plasma concentration-time profiles following intravenous administration of Fentanyl of Example 2 (n=3) in the study of Example 6.

FIG. 4 depicts the mean (±S.E.) plasma concentration-time profiles following sublingual administration of Fentanyl of 5 Example 2 (n=3) in the study of Example 6.

FIG. 5 depicts the mean (±S.E.) plasma concentration-time profiles following intravenous administration of Fentanyl of Example 3 (n=3) in the study of Example 6.

FIG. 6 depicts the mean (±S.E.) plasma concentration-time ¹⁰ profiles following sublingual administration of Fentanyl of Example 3 (n=3) in the study of Example 6.

FIG. 7 depicts the mean (±S.E.) plasma concentration-time profiles following intravenous administration of Fentanyl of Example 4 (n=3) in the study of Example 6.

FIG. 8 depicts the mean (±S.E.) plasma concentration-time profiles following sublingual administration of Fentanyl of Example 4 (n=3) in the study of Example 6.

FIG. **9** depicts the mean (±S.E.) plasma concentration-time profiles following intravenous administration of Fentanyl of ²⁰ Example 5 (n=3) in the study of Example 6.

FIG. 10 depicts the mean (±S.E.) plasma concentrationtime profiles following sublingual administration of Fentanyl of Example 5 (n=3) in the study of Example 6.

FIG. 11 depicts a graphical summary of the Dv10, Dv50, 25 Dv90, and plume records values versus placement distance for vignetting results.

FIG. 12 depicts a graphical summary of Dv10, Dv50, and Dv90 values versus placement at 4 cm device to laser beam for exhaust results.

FIG. 13 depicts a graphical summary of Dv10, Dv50, and Dv90 values versus placement at 7 cm device to laser beam for exhaust results.

FIG. **14** depicts a graphical summary of device to laser beam placement results for Dv10, Dv50, and Dv90 values ³⁵ versus distance to device to laser beam.

DETAILED DESCRIPTION OF THE INVENTION

The simplest and most prevalent administration route for 40 pharmacologic agents is by mouth. To use this method, a pharmacological agent is typically incorporated into a tablet, a capsule, or into a liquid base. Oral administration of a drug is extremely convenient, and for many drugs, it will continue to be the method of choice.

Absorption of a drug into the bloodstream after swallowing a tablet may vary from patient to patient. The absorption of the drug is typically dependent upon the movement from the stomach to the small and large intestines and the effects of secretions from these organs. Further, with the oral administration of a drug such as fentanyl to a patient, as the fentanyl enters the patient's bloodstream through the intestines and passes through the patient's liver before distribution throughout the body, upwards of fifty percent or more of the fentanyl may be removed from the patient's bloodstream. This "first 55 pass effect" results in the oral route of administration being impractical for fentanyl.

Absorption of fentanyl or a pharmaceutically acceptable salt thereof into the bloodstream following oral administration is significantly reduced by the first pass effect. Therefore, 60 the oral route of administration is impractical for fentanyl. Other forms of delivery such a parenteral, buccal, and transdermal delivery have been utilized to decrease or avoid this first pass effect for fentanyl. However, these other forms of delivery have certain disadvantages associated with them. For example, parenteral administration requires injection using a syringe and needle, and may lead to necrosis that can accom-

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pany i.m. administration of drugs; Actiq®, a transmucosal fentanyl citrate lozenge formulation requires the patient to constantly suck on the lozenge which is attached to a handle (similar to a lollipop) in order to obtain effective pain relief; and Duragesic®, a transdermal fentanyl delivery device, is suitable for the management of chronic pain, but is not indicated for acute or breakthrough pain.

The oral cavity offers a simple, painless method of opioid analgesic administration. Within the oral cavity, there are three generally recognized routes of administration of an active agent, namely local, buccal and sublingual.

Local delivery is mainly limited to applications regarding disruptions occurring within the oral cavity itself, such as a canker sore.

The buccal mucosa area encompasses the mucosal membranes of the inner lining of the cheeks. The buccal mucosa is however, less permeable than the sublingual area. One of the major disadvantages associated with buccal mucosa delivery of an active agent has been the relatively low passage of active agents across the mucosal epithelium, thereby resulting in low agent bioavailability, which translates into a substantial loss of usable active agent within each dosage.

Sublingual delivery is achieved through the mucosal membranes lining the floor of the mouth. Because of the high permeability and the rich blood supply, transport via the sublingual route results in a rapid onset of action, providing a delivery route appropriate for highly permeable drugs with short delivery period requirements and an infrequent dosing regimen.

The sublingual formulations of the present invention are useful in the treatment of moderate to severe pain. Preferably the sublingual formulations of the present invention are useful for the treatment of breakthrough pain. For example, the formulations of the present invention are preferably suitable for a patient receiving chronic pain therapy who experiences breakthrough pain and is in need of acute pain relief.

The sublingual formulations of the present invention may be used to alleviate pain from many causes, including but not limited to shock, limb amputation, severe chemical or thermal burn injury, sprains, ligament tears, fractures, wounds and other tissue injuries, dental surgery, procedures and maladies, labor and delivery, during physical therapy, post operative pain, radiation poisoning, cancer, acquired immunodeficiency syndrome (AIDS), epidural (or peridural) fibrosis, back surgery and laminectomy, sciatica, painful sickle cell crisis, arthritis, autoimmune disease, intractable bladder pain, and the like. Sublingual administration of the formulations of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, of the present invention is also preferably amenable to hospice use, particularly hospices that specialize in the care of cancer and AIDS patients.

In certain preferred embodiments, the sublingual administration of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, can relieve or alleviate episodes of acute breakthrough pain that can occur in a chronic pain condition. In a further embodiment, sublingual administration of fentanyl, pharmaceutically acceptable salt thereof, or derivative thereof can be used as an adjunct therapy to a conventional treatment regimen for a chronic pain condition to alleviate breakthrough pain. In certain embodiments, the invention also provides a formulation for use as an anesthetic premedication, for the induction of anesthesia, for use as a sedative and/or for the treatment of anxiety.

Sublingual administration of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, in accordance with the present invention may be particularly beneficial in the patient with cancer who is unable to tolerate oral admin-

istration because of nausea and vomiting, dysphagia as a result of disease, or parenteral administration because of decreased venous access, emaciation, or coagulation defects. Sublingual administration of fentanyl in accordance with the present invention preferably has potential advantages of even greater ease of use and rapid onset of pain relief action. Furthermore, because sublingual venous drainage is systemic rather than portal, hepatic first-pass elimination may be avoided. The present invention preferably provides therapeutic formulations and methods for solutions of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof to be delivered by sublingual spray pumps.

In certain preferred embodiments, the sublingual administration of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, is advantageous over other forms of administration in that it does not require injection using a syringe and needle, it avoids necrosis that can accompany i.m. administration of drugs, and it avoids the need to constantly suck on a lozenge or lollipop. Preferably the sublingual administration of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, in accordance with the present invention is suitable for self administration.

In preferred embodiments certain embodiments, the formulations of the present invention are advantageous in that 25 propellants such as hydrofluorocarbon propellants such as volatile chlorofluorocarbons (e.g. propellant 12), volatile hydrofluoroalkanes (e.g. 1,1,1,2-tetrafluoroethane and 1,1,1, 2,3,3,3-heptafluoro-n-propane) and volatile alkanes (e.g. propane, butane) are not required to deliver the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, sublingually to the patient.

Preferably the formulations of the present invention are delivered as liquid droplets having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more 35 preferably a mean diameter of from about 20 to about 200 microns. Most preferably the formulations are delivered as liquid droplets have a size distribution of from about 5 microns to about 500 microns, preferably from about 10 microns to about 200 microns, preferably from about 20 40 microns to about 100 microns, more preferably from about 30 microns to about 70 microns.

Preferably the delivery of the formulation of the present invention to the sublingual mucosa via spray results in a rapid onset of therapeutic effect of the fentanyl, a pharmaceutically 45 acceptable salt thereof, or derivative thereof. For example, in certain embodiments the formulations of the present invention provide a mean time to maximum plasma concentration (T_{max}) of fentanyl at from about 5 minutes to about 120 minutes, preferably at from about 10 to about 60 minutes, and 50 more preferably at from about 15 to about 35 minutes after sublingual administration to humans.

In certain further embodiments the formulations of the present invention provide a mean maximum plasma concentration (C_{max}) of fentanyl of about 127 pg/ml to about 213 55 pg/ml per 100 µg fentanyl, preferably about 142 pg/ml to about 195 pg/ml per 100 µg fentanyl, more preferably about 158 pg/ml to about 177 pg/ml per 100 µg fentanyl after sublingual administration to humans.

In certain further embodiments the formulations of the 60 present invention provide a mean maximum plasma concentration (C_{max}) of fentanyl of about 137 pg/ml to about 207 pg/ml based on a 100 µg fentanyl dose, preferably about 154 pg/ml to about 190 pg/ml based on a 100 µg fentanyl dose, more preferably about 163 pg/ml to about 181 pg/ml based on a 100 µg fentanyl dose after sublingual administration to humans.

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In certain further embodiments the formulations of the present invention provide a mean maximum plasma concentration (C_{max}) of fentanyl of about 566 pg/ml to about 850 pg/ml based on a 400 µg fentanyl dose, preferably about 637 pg/ml to about 779 pg/ml based on a 400 µg fentanyl dose, more preferably about 672 pg/ml to about 744 pg/ml based on a 400 µg fentanyl dose after sublingual administration to humans

In certain further embodiments the formulations of the present invention provide a mean maximum plasma concentration (C_{max}) of fentanyl of about 1016 pg/ml to about 1525 pg/ml based on a 800 µg fentanyl dose, preferably about 1143 pg/ml to about 1398 pg/ml based on a 800 µg fentanyl dose, more preferably about 1206 pg/ml to about 1334 pg/ml based on a 800 µg fentanyl dose after sublingual administration to humans

In certain further embodiments the formulations of the present invention provide an area under the plasma concentration time curve to infinity (AUC $_{\infty}$) of fentanyl of about 572 pg·h/ml to about 1273 pg·h/ml per 100 µg fentanyl, preferably about 644 pg·h/ml to about 1167 pg·h/ml per 100 µg fentanyl, more preferably about 715 pg·h/ml to about 1061 pg·h/ml per 100 µg fentanyl after sublingual administration to humans.

In certain further embodiments the formulations of the present invention provide an area under the plasma concentration time curve to infinity (AUC $_{\infty}$) of fentanyl of about 654 pg·h/ml to about 982 pg·h/ml based on a 100 µg fentanyl dose, preferably about 736 pg·h/ml to about 900 pg·h/ml based on a 100 µg fentanyl dose, more preferably about 777 pg·h/ml to about 859 pg·h/ml based on a 100 µg fentanyl dose after sublingual administration to humans.

In certain further embodiments the formulations of the present invention provide an area under the plasma concentration time curve to infinity (AUC $_{\infty}$) of fentanyl of about 3394 pg·h/ml to about 5092 pg·h/ml based on a 400 μg fentanyl dose, preferably about 3818 pg·h/ml to about 4667 pg·h/ml based on a 400 μg fentanyl dose, more preferably about 4030 pg·h/ml to about 4455 pg·h/ml based on a 400 μg fentanyl dose after sublingual administration to humans.

In certain further embodiments the formulations of the present invention provide an area under the plasma concentration time curve to infinity (AUC $_{\infty}$) of fentanyl of about 4581 pg·h/ml to about 6873 pg·h/ml based on a 800 µg fentanyl dose, preferably about 5154 pg·h/ml to about 6300 pg·h/ml based on a 800 µg fentanyl dose, more preferably about 5440 pg·h/ml to about 6014 pg·h/ml based on a 800 µg fentanyl dose after sublingual administration to humans.

In certain further embodiments the formulations of the present invention provide an area under the plasma concentration time curve from zero to the time of the last quantifiable plasma concentration (AUC $_T$) of fentanyl of about 378 pg·h/ml to about 1067 pg·h/ml per 100 µg fentanyl, preferably about 425 pg·h/ml to about 978 pg·h/ml per 100 µg fentanyl, more preferably about 472 pg·h/ml to about 889 pg·h/ml per 100 µg fentanyl after sublingual administration to humans.

In certain further embodiments the formulations of the present invention provide an area under the plasma concentration time curve from zero to the time of the last quantifiable plasma concentration (AUC $_{T}$) of fentanyl of about 378 pg·h/ml to about 568 pg·h/ml based on a 100 μg fentanyl dose, preferably about 425 pg·h/ml to about 520 pg·h/ml based on a 100 μg fentanyl dose, more preferably about 448 pg·h/ml to about 497 pg·h/ml based on a 100 μg fentanyl dose after sublingual administration to humans.

In certain further embodiments the formulations of the present invention provide an area under the plasma concentration time curve from zero to the time of the last quantifiable

plasma concentration (AUC $_T$) of fentanyl of about 2844 pg·h/ml to about 4268 pg·h/ml based on a 400 μ g fentanyl dose, preferably about 3200 pg·h/ml to about 3912 pg·h/ml based on a 400 μ g fentanyl dose, more preferably about 3378 pg·h/ml to about 3734 pg·h/ml based on a 400 μ g fentanyl dose safter sublingual administration to humans.

In certain further embodiments the formulations of the present invention provide an area under the plasma concentration time curve from zero to the time of the last quantifiable plasma concentration (AUC $_T$) of fentanyl of about 4333 pg·h/ ml to about 6501 pg·h/ml based on a 800 µg fentanyl dose, preferably about 4875 pg·h/ml to about 5960 pg·h/ml based on a 800 µg fentanyl dose, more preferably about 5146 pg·h/ ml to about 5689 pg·h/ml based on a 800 µg fentanyl dose after sublingual administration to humans.

Preferably the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, is dissolved in an organic solvent. Examples of organic solvents that may be used to enhance the solubility of fentanyl, or the pharmaceutically 20 acceptable salt thereof in a carrier such as e.g., water, include for example and without limitation: lower alcohols (e.g. C_{1-4} alcohols) such as methanol, ethanol, propyl alcohol, or butyl alcohol; C₂₋₈ alcohols having two or three hydroxyl groups, preferably glycerol, propylene glycol or butylene glycol; and 25 polyethylene glycols such as PEG_{200} and PEG_{400} and the like. Mixtures of any of the aforementioned solvents may be used. In certain embodiments, the solvent is a non-polar hydrocarbon, preferably a C₇₋₁₈ hydrocarbon of a linear or branched configuration, its alcohols, fatty acid esters, and triglycerides, such as miglyol. In certain preferred embodiments the organic solvent is ethanol, propylene glycol, polyethylene glycol, or combination thereof.

Preferably the amount of organic solvent for inclusion in the formulation is at least an amount of organic solvent necessary to adequately solubilize the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, such that the fentanyl remains in solution and does not precipitate out.

In certain embodiments, the organic solvent is included in the formulation in an amount of from about 0% to about 99.9% by weight of the formulation, preferably from about 10% to about 80% by weight of the formulation, more preferably from about 20% to about 60% by weight of the formulation.

In certain embodiments, the compositions comprise a C_{2-8} alcohol such as propylene glycol, or a polyethylene glycol and/or polypropylene glycol of an average molar weight of 200 to 4000, or a mixture thereof, in addition to the organic solvent described above. The C_{2-8} alcohol may act as a cosolvent in combination with the organic solvent. Polyethylene glycols commercially available as Carbowax® (e.g., Carbowax 300 of a molar weight of 300), can be used.

In certain embodiments, the solvent is a cosolvent which includes any of the solvents mentioned herein. In certain 55 preferred embodiments, the cosolvent includes ethanol, propylene glycol, polyethylene glycol, labrosol, labrafil, transcutol, or combination thereof.

In certain preferred embodiments, the composition according to the invention comprises from about 0.0001% to about 60 20% by weight of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof; from about 1% to about 99% by weight of organic solvent; and from about 0.01% to about 50% by weight of C_{2-8} alcohol.

In certain preferred embodiments, the composition according to the invention comprises from about 0.001% to about 15% by weight of fentanyl, a pharmaceutically acceptable

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salt thereof, or derivative thereof; from about 5% to 90% by weight of ethanol; and from about 0.1% to 40% by weight of propylene glycol.

In certain preferred embodiments, the composition according to the invention comprises from about 0.01% to about 10% by weight of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof; from about 10% to about 80% by weight of ethanol; and from about 1% to about 30% by weight of propylene glycol.

In certain preferred embodiments, the composition according to the invention comprises from about 0.1% to about 0.8% by weight of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof; from about 20% to about 60% by weight of ethanol; and from about 4% to about 6% by weight of propylene glycol.

In certain preferred embodiments, the composition according to the invention comprises in a 1 ml volume: from about 100 μg/ml to about 800 μg/ml fentanyl base, about 50% ethanol, about 5.2% propylene glycol, and water qs to 1 ml.

In certain embodiments the fentanyl is employed in the form of a pharmaceutically acceptable salt. Examples of suitable salt forms of fentanyl for use in accordance with the present invention include for example and without limitation, the hydrochloride, chloride, sulphate, tartrate, or citrate salt forms. In certain preferred embodiments, the fentanyl is employed as the free base in the formulations of the present invention.

In certain preferred embodiments, the fentanyl, pharmaceutically acceptable salt thereof, or derivative thereof, will be employed in the formulation at a concentration of from about 0.05 mg/ml to about 15 mg/ml, preferably from about 0.1 mg/ml to about 10 mg/ml, more preferably from about 1 mg/ml to about 8 mg/ml (where weight is expressed as weight of fentanyl free base).

In certain preferred embodiments, the amount of fentanyl, pharmaceutically acceptable salt thereof, or derivative thereof; delivered per unit dose is about $10 \,\mu g$ to about $10 \,mg$, preferably from about 25 μg to about 5 mg, more preferably from about 50 μg to about 1600 μg .

In preferred embodiments of the present invention, the formulation is a solution. In certain alternate embodiments, the formulation is a suspension. When the formulation of the present invention is a suspension, it may be necessary to shake the composition prior to spraying.

In certain preferred embodiments, after the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, is dissolved in the organic solvent, the formulation is preferably included into a liquid carrier for the delivery of the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, via a spray device.

Pharmaceutically acceptable carriers include but are not limited to water, buffer, saline, buffered saline, dextrose solution, propylene glycol, polyethylene glycols, miglyol, and the like. In a specific embodiment, a carrier that may be used in the pharmaceutical formulation of the present invention is phosphate buffered saline, or a buffered saline. In certain preferred embodiments the carrier is water. In certain embodiments, the water in the formulation is present in the form of an aqueous buffer. The buffer is preferably adapted to stabilize the pH of the formulation at pH of about 5 to about 12, preferably at pH of about 6 to about 10, more preferably from about 8 to about 9.5. Buffer systems for use in accordance with the present invention include for example and without limitation sodium acetate/acetic acid, ammonium acetate/disodium edentate, boric acid/sodium hydroxide,

orthophosphoric acid/sodium hydroxide, sodium hydrogen carbonate/sodium carbonate, disodium hydrogen orthophosphate/citric acid, and the like.

Other components such as preservatives, antioxidants, surfactants, absorption enhancers, viscosity enhancers or film forming polymers, bulking agents, diluents, coloring agents, flavoring agents, pH modifiers, sweeteners or taste-masking agents may also be incorporated into the composition. Suitable coloring agents include red, black and yellow iron oxides and FD&C dyes such as FD&C Blue No. 2, FD&C Red No. 40, and the like. Suitable flavoring agents include mint, raspberry, licorice, orange, lemon, grapefruit, caramel, vanilla, cherry grape flavors, combinations thereof, and the like. Suitable pH modifiers include citric acid, tartaric acid, phosphoric acid, hydrochloric acid, maleic acid, sodium hydroxide, and the like. Suitable sweeteners include aspartame, acesulfame K, thaumatic, and the like. Suitable taste-masking agents include sodium bicarbonate, ion-exchange resins, cyclodextrin inclusion compounds, adsorbates, and the like.

Absorption enhancers for use in accordance with the present invention include, for example, polysorbates, sorbitan esters, poloxamer block copolymers, PEG-35 castor oil, PEG-40 hydrogenated castor oil, caprylocaproyl macrogol-8 glycerides, PEG-8 caprylic/capric glycerides, sodium lauryl 25 sulfate, dioctyl sulfosuccinate, polyethylene lauryl ether, ethoxydiglycol, propylene glycol mono-di-caprylate, glycerol monocaprylate, glyceryl fatty acids (C8-C18) ethoxylated, oleic acid, linoleic acid, glyceryl caprylate/caprate, glyceryl monooleate, glyceryl monolaurate, caprylic/capric 30 triglycerides, ethoxylated nonylphenols, PEG-(8-50) stearates, olive oil PEG-6 esters, triolein PEG-6 esters, lecithin, d-alpha tocopheryl polyethylene glycol 1000 succinate, polycarbonate, sodium glycocholate, sodium taurocholate, cyclodextrins, citric acid, sodium citrate, triacetin, combinations 35 thereof, and the like. In certain preferred embodiments, the absorption enhancer is triacetin. In certain preferred embodiments wherein an absorption enhancer is included in the formulation, the absorption enhancer is included in an amount of from about 0.001% to about 10% by weight of the 40 formulation, preferably in an amount of about 0.01% to about 5% by weight of the formulation.

Bulking agents for use in accordance with the present invention include for example, microcrystalline cellulose, mannitol, xylitol, starches and the like. In certain preferred 45 embodiments, the bulking agent is mannitol. In certain preferred embodiments wherein bulking agent is included in the formulation, the bulking agent is included in an amount of from about 0.001% to about 10% by weight of the formulation, preferably in an amount of about 0.01% to about 5% by 50 weight of the formulation.

Film-forming polymers for use in accordance with the present invention may serve for decreasing the fineness of the spray, the spraying angle and preferably the spreading by increasing the viscosity of the composition. As film-forming 55 polymer, gellan gum, xantham gum, carboxymethyl cellulose, hydroxypropyl cellulose, hydroxyethyl cellulose, hydroxypropyl methylcellulose, methylcellulose, ethylcellulose, gelucire, poloxamers, alginic acid, propyleneglycol ester, polyvinyl alcohol (PVA), polyvinylpyrrolidone (PVP), 60 PVP/PVA copolymer, lubrajel, carboxyvinyl polymer, acrylic acid polymers and copolymers, methacrylic acid, methyl acrylate, ethyl acrylate, methyl methacrylate, ethyl methacrylate, combinations thereof and the like can be used. In certain embodiments, an increase in the viscosity of the 65 solution using film-forming polymers or the like provides an increase in the droplet size when administered from the spray

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device. The chemistry of the polymer and the molecular weight of the polymer may also influence the diameter of the droplets.

In certain embodiments, the formulations according to the invention are preferably packaged as a bulk solution containing multiple doses in a pump spray system comprising a sealed container fitted with a metering pump.

In certain alternate embodiments the formulations according to the invention are preferably package as a single unit dose solution in a single unit dose pump spray system comprising a sealed container fitted with a pump.

Typically a patient is treated by administration sublingually of 1 to 2 actuations, from the spray pump. Another advantage of sublingual spray delivery is the ability to easily titrate patients by 1 or 2 doses as required by a single actuation. This is typically not the case with other forms of drug delivery (patches, lozenges, tablets, suppositories).

Pump action sprays are characterized in requiring the application of external pressure for actuation, for example, external manual, mechanical or electrically initiated pressure. This is in contrast to pressurized systems, e.g., propellant-driven aerosol sprays, where actuation is typically achieved by controlled release of pressure e.g., by controlled opening of a valve.

In certain embodiments the pump sprays are preferred as the use of a pump spray with the formulation of the present invention allows for the administration of droplets or particles having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more preferably a mean diameter of from about 20 to about 200 microns, and/or preferably having a size distribution of from about 5 microns to about 500 microns, preferably from about 10 microns to about 200 microns, preferably from about 20 microns to about 100 microns, more preferably from about 30 microns to about 70 microns. This is in contrast to a pressurized system which may result in particles less than 5 microns. Liquid droplets or particles having a diameter of less than about 5 microns have the potential to enter into the lungs of a human upon administration. Such entry into the lungs could lead to an increase in patient to patient variability in absorption of the fentanyl. Further, absorption of fentanyl in the lungs could lead to an increased absorption and increased side effects, including respiratory depression which may be fatal.

In certain preferred embodiments, the droplet size of the delivered formulations further provides for an increase in surface area by being sprayed sublingually as opposed to being placed under the tongue with e.g., a dropper.

In certain preferred embodiments, the delivery device is a device such as those described in U.S. Pat. Nos. 6,866,566; 6,877,672; 6,772,915; 6,725,857; 6,705,493; 6,679,248; 6,578,741; 6,527,144; 6,484,715; 6,478,196; 6,461,322; 6,446,839; 6,427,878; 6,367,473; 6,364,166; 6,321,942; 6,234,366; 6,227,413; 6,059,151; 6,059,150; 6,055,979; 5,944,222; 5,901,883; 5,813,570; 4,565,302; 4,532,967; 6,964,381; 6,860,411; 6,824,020; 6,817,490; 6,585,172; 6,443,370; 6,427,680; 6,425,499; 6,401,987; 6,398,074; 6,264,065; 5,950,877; 5,328,099; 5,301,846; and the like which are described in certain embodiments as being suitable for nasal administration.

Other devices suitable for use in accordance with the formulations of the present invention are described in U.S. Pat. Nos. 6,808,085; 6,736,293; 6,732,955; 6,708,846; 6,626,379; 6,626,330; 6,626,328; 6,454,185; 6,427,876; 6,427,684; 6,419,167; 6,405,903; 6,352,181; 6,308,867; 6,257,461; 6,257,454; 6,250,509; 6,227,415; 6,209,760; 6,179,164; 6,109,547; 6,062,430; 6,026,992; 5,992,704; 5,992,703; 5,988,449; 5,967,369; 5,964,417; 5,950,879; 5,938,125;

5,927,559; 5,921,444; 5,893,484; 5,875,938; 5,862,962; 5,860,567; 5,816,504; 5,813,570; 5,803,311; 5,791,518, 5,692,650; 5,655,689; 5,584,417; 5,520,337; 5,519,980; 5,482,193; 5,469,989; 5,443,185; 5,439,177; 5,437,398; 5,427,280; 5,395,032; 5,375,745; 5,368,201; 5,366,122; 5,366,122; 5,335,823; 5,326,000; 5,323,936; 5,316,198; 5,301,841; 5,295,628; 5,289,946; 5,277,334; 5,257,726; 5,228,586; 5,209,375; 5,203,840; 5,147,087; 5,115,980; 5,110,052; 5,011,046; 4,958,752; 4,946,069; 4,944,430; 4,934,568; 4,921,142; 4,871,092; 4,830,284; 4,826,048; 4,823,991; 4,821,923; 4,817,829; 4,776,498; 4,762,475; 4,728,008; 4,726,747; 4,694,977; 4,694,976; 4,566,611; 6,851,583; 6,824,021; 6,779,690; 6,776,312; 6,971,559; 6,948,640; 6,945,473; 6,938,802; 6,933,850; 6,929,156; 6,918,514; 6,913,205; 6,866,168; 6,832,072; 6,830,163; 15 6,817,490; 6,817,489; 6,811,060; 6,811,057; 6,805,301; 6,805,263; 6,789,750; 6,789,706; 6,786,369; 6,783,035; 6,772,913; 6,769,579; 6,758,371; 6,752,298; 6,742,677; 6,705,062; 6,698,627; 6,698,623; 6,663,019; 6,659,314; 6.659,307; 6.655,550; 6.655,549; 6.651,846; 6.601,735; 20 6,595,395; 6,592,010; 6,588,629; 6,581,852; 6,571,991; 6,554,160; 6,536,635; 6,527,149; 6,527,148; 6,488,185; 6,471,097; 6,460,781; 6,460,740; 6,460,738; 6,446,841; 6,422,429; 6,409,049; 6,398,079; 6,360,919; 6,349,856; 6,345,737; 6,343,722; 6,662,561; 6,315,169; 6,273,303; 25 6,273,300; 6,261,274; 6,257,457; 6,234,363; 6,234,168; 6,221,054; 6,209,759; 6,189,741; 6,186,371; 6,155,496; 6,119,897; 6,105,826; 6,021,930; 6,012,615; 5,988,496; 5,950,871; 5,931,386; 5,850,948; 5,803,318; 5,799,810; 5,769,325; RE35,683; 5,692,492; 5,568,884; 5,566,865; 30 5,511,698; 5,482,188; 5,476,198; 5,366,115; 5,337,923; 5,249,713; 5,237,797; 5,234,135; 5,226,563; 5,190,192; 5,176,296; 5,127,548; 4,966,313; 4,91,840; 4,245,967; 4,030,667; and the like.

All of the patents recited herein are hereby incorporated by 35 reference in their entireties. Although the delivery devices disclosed in the patents described above may be suitable for nasal or inhalation administration, in accordance with certain embodiments of the present invention the delivery devices are specifically adapted to be suitable for sublingual administration of a liquid formulation. In certain embodiments, the devices utilized to practice the present invention include components made from Pfeiffer of America, Inc., for example, Pfeiffer of America, Inc. sublingual unit dose device article reference number 72772. In other embodiments, the device is 45 Pfeiffer of America, Inc., sublingual unit dose applicator assembly.

Preferably the device in accordance with the present invention is adapted to sublingually deliver the sublingual formulation in a controlled manner preferably such that only droplets having a mean diameter of at least about 10 microns, preferably at least about 20 microns, more preferably a mean diameter of from about 20 to about 200 microns are delivered to the patient. More preferably only droplets having a size distribution in the range of from about 5 microns to about 500 smicrons, preferably from about 10 microns to about 200 microns, preferably from about 20 microns to about 100 microns, more preferably from about 30 microns to about 70 microns.

Preferably the dispenser is constructed in such a way that it 60 can be carried and simultaneously reliably operated with the fingers, or with three fingers of one hand, and can be used, for example, in the manner of a sublingual spray. The dispenser can be constructed as a disposable dispenser which, following the emptying of the medium chamber, does not have to be 65 refilled and can therefore be constructed as a simple standard component, which receives the pump, the formulation, the

channels and optionally, valves or closures within an outer casing, which in side view can be roughly T-shaped or Y-shaped.

If the dispenser is to be emptied in a single pump stroke in successive portions or in one complete pump stroke, and is not to be refilled, then the dispenser can be substantially tightly closed with respect to the outside in the starting position

In certain preferred embodiments, the delivery device (e.g., such as a spray pump device) includes a lock-out mechanism. Preferably the lock-out mechanism allows for administration of only one unit dose, and preferably prevents abuse of the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, by only allowing for the administration of one dose and locking out of further administration for a certain and/or predetermined period of time. In certain embodiments, after one or more actuating cycles the actuator can be automatically transferred into the locking position, so that for performing a following actuating cycle randomly or deliberately a release must take place. Locking can take place in the starting position, actuating position and/or an intermediate position and can act both against actuation and against return or against one of these movements alone and several locking positions with the same or different locking action are pos-

In certain embodiments, the device may be premetered or alternatively, the device may be device-metered. Premetered devices preferably contain previously measured doses or a dose fraction in some type of units (e.g., single unit dose amount of solution, single or multiple blisters or other cavities) that may be included in the device during manufacture or by the patient before use. Typical device-metered units have a reservoir containing formulation sufficient for multiple doses that are delivered as metered sprays by the device itself when activated by the patient.

Important factors to consider with manufacture of the device are the reproducibility of the dose, the spray plume, and the particle/droplet size distribution, which can affect the delivery of the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, under the tongue. Maintaining the reproducibility of these parameters through the expiration dating period and ensuring the functionality of the device (e.g., spray mechanism, electronic features, sensors) through its lifetime under patient-use conditions is important as any alteration in these parameters could lead to variability in dosing and absorption, which could lead to potential side effects.

The administered dose of spray drug formulation may be dependent on the design, reproducibility, and performance characteristics of the container closure system. A suitable device which provides the desired droplet/particle size distribution is an important factor for the correct performance of the fentanyl product. Actuation parameters (e.g., force, speed, hold and return times) should also be considered with respect to the device. Moreover, the device should be compatible with formulation components. Further, the device should be designed to prevent partial metering of the fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, formulation when used according to the patient instructions for use.

A typical device includes a base unit, a discharge actuator, an orifice for the formulation to be release from the device, and a medium reservoir. Preferably a reservoir is provided which as a dispensing chamber is filled already on production of the device. The medium reservoir preferably defines a

measured content of fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, to be discharged upon a single activation.

In accordance with certain embodiments of the invention, a reservoir, or a space thereof receiving the medium is preferably an elongated shape preferably having a wall thickness which is constant over the circumference and length of the reservoir body. The reservoir body may be formed simply by a section of a cylindrical hollow of a plastics, steel, such as stainless steel, transparent material, such as glass, or the like so that its production is very simple.

Preferably an actuator body is provided on a unit of the device, which is movable relative to the orifice for activating discharge. This body, in the course of the actuating movement, opens a closure of a chamber, e.g. by puncturing. The space within this chamber may directly adjoin the medium in the reservoir, accommodate the opening body or the reservoir at least in part and configured as a pressure space which prior to being opened is at an elevated pressure. The opening body may be formed directly by the reservoir.

Preferably during a part of the actuating travel following the starting position an elevated pressure is built up. In a subsequent portion of the actuating movement continuing in the same direction, the medium is relieved of the pressure at 25 one of the sides and communicated to the medium orifice on this side. As such, due to the pressure acting on the side, the medium is pushed from the reservoir and through the orifice.

Typically as the liquid formulation leaves the orifice, the liquid droplets follow a trajectory which is influenced by the orifice shape of the device. In certain embodiments, the droplet size, spray geometry, and the spray pattern are dependent on the design of the pump and/or the properties of the formulation. In certain embodiments, the orientation of the actuator, pump design, and the properties of the formulation will influence the spray symmetry and the shape.

In certain embodiments, the device of the present invention further includes a stopper. Preferably the stopper comprises a material which precludes or substantially precludes the 40 absorption of the fentanyl, pharmaceutically acceptable salt thereof, or derivative thereof. A suitable stopper for use in accordance with the device of the present invention is, for example, a stopper marketed by West Pharmaceutical Services, Inc. In certain preferred embodiments, the stopper has 45 the following composition and characteristic: 1) elastomer: bromobutyl and/or chlorobutyl; 2) reinforcement: inert material; and 3) curing system: unconventional.

In certain embodiments, the device further includes a gasket. Preferably the gasket comprises a material which precludes or substantially precludes the absorption of the fentanyl, pharmaceutically acceptable salt thereof, or derivative thereof. A suitable gasket for use in accordance with the device of the present invention is, for example, a stopper marketed by West Pharmaceutical Services, Inc. In certain 55 preferred embodiments, the gasket has the following composition and characteristic: 1) elastomer: bromobutyl and/or chlorobutyl; 2) reinforcement: inert material; and 3) curing system: unconventional.

Droplet size distribution can be determined by utilizing any 60 reliable method known to one of skill in the art. One such method uses laser diffraction devices, such as, for example, the Malvern Spraytec® with RT Sizer software. A Malvern Mastersizer S, by Malvern Instruments Limited (U.K.), device may also be used to determine size distribution. A 65 Malvern Mastersizer S is a modular particle size analyzer offering measurement versatility. It can measure spray drop-

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let size as well as wet and dry samples. Particles from submicron to a few millimeters may be measured with the Malvern Mastersizer S.

Further, automated actuation stations for comparative in vitro bioequivalence tests or other testing to decrease the variability associated with manual actuation may also be used when determining the droplet size distribution. Any such automated actuation stations known to one of skill may be applicable in practicing the present invention. An example of one such device is the MightyRunt Actuation Station by Innova Systems, Inc. In a preferred embodiment, a MightyRunt is equipped with an exhaust fan attachment. In a further embodiment, the MightyRunt is further equipped with a Mettler Toledo balance Model AT201.

5 Other Active Agents

Although the invention described herein has been described with respect to fentanyl, a pharmaceutically acceptable salt thereof, or derivative thereof, it is contemplated that other active pharmaceutical agents, particularly those suitable for sublingual administration, may be used in accordance with the present invention. For example, in certain embodiments, other opioid analgesics which are suitable for sublingual administration may be used in place of the fentanyl, pharmaceutically acceptable salt thereof, or derivative thereof. Certain opioid analgesics for use in accordance with the present invention include, for example and without limitation, alfentanil, sufentanil, buprenorphine, butorphanol, codeine, hydrocodone, hydromorphone, levorphanol, meperidine, methadone, morphine, nalbuphine, oxycodone, oxymorphone, propoxyphene, tramadol, and the like.

In certain embodiments the active compound is an analgesic, opioid antagonist, anti-migraine agent, anti-emetic agent, anti-epileptic agent, anti-hypertensive agent, anesthetic agent, cannabinoid, cannabinoid antagonist, inverse agonist of cannabinoid, endocannabinoid, enkephalin, analogues or derivatives thereof, or mixtures thereof.

Analgesics useful in the present invention include, but are not limited to, alfentanil, buprenorphine, butorphanol, codeine, fenpipramide, hydrocodone, hydromorphone, levorphanol, meperidine, methadone, morphine, nalbuphine, oxycodone, oxymorphone, pentazocine, piritramide, propoxyphene, sufentanil, tilidine, tramadol, analogues or derivatives thereof, or mixtures thereof.

Opioid antagonists useful in the present invention include, but are not limited to, naloxone, naltrexone, nalmefene, analogues or derivatives thereof, or mixtures thereof.

Anti-migraine agents useful in the present invention include, but are not limited to, almotriptan, eletriptan, frovatriptan, naratriptan, rizatriptan, sumatriptan, zolmitriptan, ergot alkaloids, proxibarbal, lisuride, methysergide, clonidine, pizotifene, analogues or derivatives thereof, and mixtures thereof.

Anti-emetic agents useful in the present invention include, but are not limited to bromopride, domperidone, granisetron, ondansetron, tropisetron, metoclopramide, pyridoxine, scopolamine, thiethylperazine, analogues or derivatives thereof, or mixtures thereof.

Anti-epileptic agents useful in the present invention include, but are not limited to barbiturates, carbamazepine, ethosuximide, mesuximide, phenyloin, primidone, sultiam, valproic acid, vigabatrine, analogues or derivatives thereof, or mixtures thereof.

Anti-hypertensive agents useful in the present invention include, but are not limited to diltiazem, clonidine, nifedipine, verapamil, isosorbide-5-mononitrate, organic nitrates, agents used in treatment of heart disorders, analogues or derivatives thereof, or mixtures thereof.

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Anesthetics useful in the present invention include, but are not limited to benzocaine, bupivacaine, dibucaine, etidocaine, levobupivacaine, lidocaine, mepivacaine, oxybuprocaine, piperocaine, prilocaine, procaine, proparacaine, ropivacaine, tetracaine, xylocaine, desflurane, enflurane, isoflurane, sevoflurane, benzonatate, dyclonine, ketamine, phenol, propofol, analogues or derivatives thereof, or mixtures thereof.

Cannabinoids useful in the present invention include, but are not limited to delta-8-tetrahydrocannabinol, delta-9-tetrahydrocannabinol, cannabidol, olivetol, cannabinol, cannabigerol, nabilone, delta-9-tetrahydro cannabinotic acid, the non-psychotropic cannabinoid 3-dimethylnepty 11 carboxylic acid homologine 8, delta-8-tetrahydrocannabinol, pharmaceutically acceptable salts thereof, complexes thereof, derivatives thereof, or mixtures thereof. A particularly preferred cannabinoid is delta-9-tetrahydrocannabinol, also known as dronabinol.

Further, active agents having a narrow therapeutic index or range (e.g., wherein small variances in blood levels of the drug causes changes in the effectiveness or toxicity of that drug) could be particularly suitable for use in accordance with the present invention. Such active agents include for example and without limitation, digoxin, levothyroxine, aminoglycosides (e.g., gentamycin, tobramycin), antiarrythimics (e.g., procainamide, quinidine), theophylline, antineoplastics, busulfan, methotrexate, 6-MP, carboplatinum, antidepressants (e.g., lithium), anticonvulsants (e.g., phenyloin, carbamazepine, valproate sodium, valproic acid), antipsychotics, anticoagulants (e.g., warfarin), cyclosporine, and the like.

The present invention will now be more fully described with reference to the accompanying examples. It should be understood, however, that the following description is illustrative only and should not be taken in any way as a restriction on the generality of the invention specified above.

DETAILED DESCRIPTION OF PREFERRED EMBODIMENTS

EXAMPLE 1

In Example 1, a fentanyl sublingual formulation was prepared having a concentration of 0.5 mg/ml. The formulation is listed in Table 1 below:

TABLE 1

Ingredient	Percent
Concentration of Fentanyl Base	percent to make 0.5 mg/m
Ethanol % (v)	20
Propylene glycol % (v)	5
DI Water % (v)	QS

EXAMPLE 2

In Example 2, a fentanyl sublingual formulation was prepared having a concentration of 0.5 mg/ml. The formulation is listed in Table 2 below:

TABLE 2

Ingredient	Percent	
Concentration of Fentanyl	percent to make 0.5 mg/ml	6

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TABLE 2-continued

	Ingredient	Percent	
5	Ethanol % (v) Propylene glycol % (v) DI Water % (v)	20 5 QS	

Contains fentanyl citrate equivalent to 0.5 mg/ml of fentanyl base

EXAMPLE 3

In Example 3, a fentanyl sublingual formulation was prepared having a concentration of 0.5 mg/ml. The formulation is listed in Table 3 below:

TABLE 3

	Ingredient	Percent
20	Concentration of Fentanyl Base	percent to make 0.5 mg/ml
	Ethanol % (v)	20
	Propylene glycol % (v)	5
	DI Water % (v)	QS
	Mannitol % (wt)	0.3
	Tween 80% (wt)	0.2
15	Tween 80% (wt)	0.2

EXAMPLE 4

In Example 4, a fentanyl sublingual formulation was prepared having a concentration of 0.5 mg/ml. The formulation is listed in Table 4 below:

TABLE 4

Ingredient	Percent
Triacetin % (wt)	0.5
Concentration of Fentanyl	percent to make 0.5 mg/ml
Base	
Ethanol % (v)	20
Propylene glycol % (v)	5
Buffer % (v)	QS

EXAMPLE 5

In Example 5, a fentanyl sublingual formulation was prepared having a concentration of 0.5 mg/ml. The formulation is listed in Table 5 below:

TABLE 5

Ingredient	Percent
Concentration of Fentanyl Base Ethanol % (v) Propylene glycol % (v) Miglyol % (v)	percent to make 0.5 mg/ml QS 5 50%

Preparation of Formulations (Examples 1-5)

- Calculated amount of Fentanyl base or Fentanyl citrate was weighed in a tared glass container.
- Calculated amount of alcohol was added to the container and mixed to dissolve fentanyl.
- Propylene glycol was weighed and added to the fentanyl solution.

- 4. Water or Buffer or Miglyol was weighed, added to the fentanyl solution and mixed for 2 mm.
- Inactive ingredients (Mannitol, Triacetin, or TW80) were added at the end and mixed well.
- The final solution was vortexed for 3 min. After mixing, the formulations were stored in refrigerator for further studies.

EXAMPLE 6

In Example 6, three rabbits weighing 2-3 kg were used to study the bioavailability of sublingual administration of Examples 1-5 in comparison to the IV injection of the formulation as a reference. Rabbits were first anesthetized by Isoflurane gas as needed to keep the rabbits immobilized for approximately 15-20 min.

For each formulation, study rabbits received a single dose of 0.1 ml (equivalent to $50\,\mu g$ of fentanyl base) by sublingual and IV administration. For the sublingual studies, the dose of liquid formulation was administered underneath the tongue using a spray bottle. Blood samples (1 ml per sample) were obtained through a catheter installed inside the ear vein.

Blood samples were collected at zero time baselines and at 5, 10, 20, 45, 60 and 120 min after the single dose. Samples were immediately cooled and plasma was separated by centrifugation within 2-3 hrs of blood collection. Samples were stored at -20° C. until assayed. After recovering from anesthesia, animals were returned to their cages. These animals were rested in cages for at least 5-7 days before they could be reused for further testing.

Plasma Collection & Separation

Blood was collected from rabbits in 3 ml tubes containing 7.5% EDTA and centrifuged at 3,000 RPM for 15 min to remove blood cells and other insoluble material. The plasma was decanted into a silanized tube and kept frozen at -20° C. until assay.

Plasma Extraction

For each plasma sample (0.5 ml), $100 \,\mu$ l of Sufentanil (IS) and $100 \,\mu$ L 5M NaOH (for protein denaturation) were added. The Fentanyl was extracted with 1 ml of 1-chlorobutane by vortex mixing for 15 min. After centrifuging at 12000 RPM for 5 min and freeze-drying for 10 min (to break up any emulsion), the upper organic layer was decanted and evaporated to dryness using a gentle stream of nitrogen. The extraction residue was reconstituted with $100 \,\mu$ l methanol followed by vortex mixing for 5 min and sonicating for 3 min, then a 3 μ l volume was injected into the GC-MS system.

Results and Discussion

Plasma Concentration vs Time Profiles

Plasma concentration-time profiles of Fentanyl in rabbits following IV and sublingual administration of $0.1\,\mathrm{ml}$ (equivalent to $50\,\mu\mathrm{g}$ of fentanyl base) are given in the following tables (Tables $6\mathrm{A}\text{-}10\mathrm{B}$).

Tables 6A and 6B provide the plasma concentration-time 55 profiles for Example 1.

TABLE 6A

(Example 1 intravenous)							
TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV		
		785.8012	839.9696	812.885396	38.302813		
10		507.7546	715.6065	611.680527	146.97351		
20	737.3449	423.5903	667.9939	609.643027	164.81539		
45	701.5631	482.215	382.5801	522.119422	163.19256		

22 TABLE 6A-continued

(Example 1 intravenous)							
TIME	R1	R2	R3	GRAND	STD DEV		
(min)	(ng/ml)	(ng/ml)	(ng/ml)	AVG			
60	562.501	554.8479	475.4686	530.939148	48.191101		
90	207.5541	492.4037	429.1217	376.359829	149.57498		
120	214.0297	492.4037	196.9675	205.4986	12.064797		

TABLE 6B

		(Example 1 sublingual)				
	TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV
	5		412.1095	370.7992	391.454361	29.210825
	10	205.5274	279.3298	755.3469	413.401352	52.186202
	20	539.2677	627.0507	712.6511	626.323191	86.693973
	45		618.9493	387.7606	503.35497	163.47506
	60	349.3563	218.14	498.3773	355.291188	140.2129
ı	90	245.8519	249.5091	231.7688	242.376606	9.3668848
	120	214.1339	162.9939	199.6146	192.247465	26.353936

In Table 7A and 7B are the plasma concentration-time profiles for Example 2.

TABLE 7A

	(Example 2 intravenous)									
TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV					
5	671.5152	788.2252		729.870183	58.35497					
10	379.1866	617.5517	1102	699.579446	300.73378					
20	603.9696	407.5375		505.75355	98.216024					
45	380.9878		292.998	336.992901	43.994929					
60	140.7566	266.3611	388.5314	265.216362	101.1569					
90	81.06491	254.4848	334.8012	223.450304	105.88636					
120	95.34888	278.7789	232.4037	202.177147	77.875432					

TABLE 7B

	(Example 2 sublingual)									
TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV					
5	350.2297	106.9249	337.6856	264.946755	136.99455					
10	373.5822	115.1643	486.0254	324.923935	190.15835					
20	285.5994	294.5517	518.5091	366.220081	131.96212					
45	302.7099	52.78093	359.9483	238.479716	163.34652					
60	118.3915	43.02434	314.9564	158.790737	140.39528					
90	81.06491	43.39148		62.2281947	26.639136					
120	95.34888	35.34888	30.68154	53.7931034	36.063946					

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In Tables 8A and 8B are the plasma concentration-time profiles for Example 3.

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TABLE 8A

	(Example 3 intravenous)									
TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV					
5	903.4949	1000.205	882.0649	928.588235	51.390778					
10	814.3834	557.9432	484.5233	618.949966	141.40552					
20		348.8641	309.1785	329.021298	19.842799					
45	120.2677	169.7972	211.0669	167.043949	37.119701					
60	160.3347	121.5882	128.9087	136.943881	16.807631					
90	89.85598	85.70081	92.71197	89.4229209	2.8786265					
120										

TABLE 8B

TABLE 9B-continued

		(Examp	le 3 sublingu	ial)					(Examp	le 4 sublingu	ıal)	
TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV		TIME	R1	R2	R3	GRAND	
5		127.3124	269.1927	198.252535	100.32454	20	(min)	(ng/ml)	(ng/ml)	(ng/ml)	AVG	STD DEV
10	304.6288	589.1684	310.5953	401.464165	162.58397							
20	349.3611	689.7999	281.5365	440.232477	218.77603		45	538.3245	499.217	449.574	495.705206	44.479353
45	288.0639	418.1555	195.9391	300.719518	111.64744		60	367.4361	452.9817	442.7566	421.058147	46.718636
60	173.0345	255.9016		214.45	74.967507		90	309.5538	571.0487	202.8722	361.158215	189.43533
90	224.432	87.7931	228.6156	180.280257	80.123534	25						
120	96.5284	145.1907		120.859533	34.409422		120	205.6531	195.7282	427.1927	276.191346	130.86513

In Tables 9A and 9B are the plasma concentration-time profiles for Example 4.

In Tables 10A and 10B are the plasma concentration-time profiles for Example 5.

TABLE 10A

	(Example 5 intravenous)									
TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV					
5	928.5193	1247.247	1123.335	1099.70047	160.67311					
10	920.1521	1100.3	1103.844	1041.43205	105.04643					
20	876.3793	966.998	972.8114	938.729547	54.075067					
45	765.9696	938.3834	947.0609	883.804598	102.14032					
60	645.6045	892.6836	482.2799	673.52265	206.62129					
90	157.9533	418.6034	343.0811	306.545977	134.1109					
120	58.43813	30.68154	30.68154	39.933739	16.025276					

TABLE 9A

Example 4 intravenous)									
TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV				
5	740.3759	732.5855	717.8438	730.268425	11.443374	50			
10	666.2644	671.4571	627.0568	654.926076	24.274753				
20	713.355	592.5	638.7444	648.199797	60.979784				
45	575.2667	557.7789	482.8316	538.625761	49.103907				
60	409.8756	596.6278	548.1217	518.208362	96.903054				
90	455.8567	484.928	430.5538	457.112801	27.208875				
120	385.2982		452.6998	418.998986	47.660144	55			

TABLE 9B

	(Example 4 sublingual)									
TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV					
5	250.8854	481.6146	459.2779	397.259297	127.25455					
10	478.074	577.9594	614.8479	556.960446	70.76361					
20	587.8195	465.9229	518.4341	524.058824	61.142626					

TABLE 10B

	(Example 5 sublingual)									
	TIME (min)	R1 (ng/ml)	R2 (ng/ml)	R3 (ng/ml)	GRAND AVG	STD DEV				
50	5	140.7911	176.7338	127.357	148,293949	25.529127				
	10	239.712	191.2956	210.8458	213.951149	24.357082				
	20	409.5335	580.6095	540.5842	510.242394	89.483087				
	45	351.3955	500.9315	314.7343	389.020453	98.636103				
5.5	60	364.9493	439.2885	280.645	361.62762	79.373899				
,,	90	245.2698	403.4255	30.68154	226.458925	187.08258				
	120	30.68154	119.7079	215.6552	122.014875	92.508392				

Pharmacokinetic Parameters

The pharmacokinetic parameters Peak plasma concentration (Cmax), Time to reach Cmax (Tmax), Half-life (t_{1/2}), Area under the curve (AUC), and Total body clearance (CL) obtained after sublingual or IV drug administrations are tabulated in the following tables (Tables 11-15). Also, plasma concentration-time curves after administering IV and sublingual doses of Fentanyl are shown in FIGS. 1-10.

TABLE 11

(Example 1 intravenous and sublingual)								
ROUTE OF ADMINISTRATION	AUC (min * ng/ml)	T _{1/2} (min)	TMAX (min)	CMAX (ng/ml)	VD (ml)	Cl (ml/min)		
IV SL	71772.86 59684.34	43.8141 55.704	0 20	812.8854 626.3232	50.7843	0.6966		

TABLE 12

(Example 2 intravenous and sublingual)									
ROUTE OF ADMINISTRATION	AUC (min * ng/ml)	T _{1/2} (min)	TMAX (min)	CMAX (ng/ml)	VD (ml)	Cl (ml/min)			
IV SL	86790.18 23759.14	153.2372 33.1477	0 20	729.8702 366.2201	181.8356	0.9218			

TABLE 13

(Example 3 intravenous and sublingual)									
ROUTE OF ADMINISTRATION	AUC (min * ng/ml)	T _{1/2} (min)	TMAX (min)	CMAX (ng/ml)	VD (ml)	Cl (ml/min)			
IV SL	32707.76 40623.13	49.7523 57.6038	0 20	928.5882 440.2325	78.4776	1.5287			

TABLE 14

(Example 4 intravenous and sublingual)									
ROUTE OF ADMINISTRATION	AUC (min * ng/ml)	T _{1/2} (min)	TMAX (min)	CMAX (ng/ml)	VD (ml)	Cl (ml/min)			
IV SL	185485.6 86916.91	200.3556 93.4018	0 10	730.2684 556.9604	77.036	0.2696			

TABLE 15

(Example 5 intravenous and sublingual)										
ROUTE OF ADMINISTRATION	AUC (min * ng/ml)	T _{1/2} (min)	TMAX (min)	CMAX (ng/ml)	VD (ml)	Cl (ml/min)				
IV SL	76250.66 42554.59	17.0944 38.2788	0 20	1099.7005 510.2424	27.3544	0.6557				

The maximum concentrations were reached in about 20 minutes for all formulations after sublingual administration. There was considerable inter-individual variability by both routes of administration. Measurable plasma concentration after sublingual administration was demonstrable after 120 min in most formulation testing. In conclusion, sublingual fentanyl administration showed good absorption profile compared to IV.

EXAMPLE 7

In Example 7, a study was conducted to determine the pharmacokinetics of a formulation prepared in accordance

with Example 1 after increasing sublingual dose administration in healthy volunteers under fasting conditions. The study was also conducted to determine the safety and tolerability of a fentanyl sublingual spray prepared in accordance with 60 Example 1 in humans.

The study was a single center, single dose, single-blinded, sequential ascending dose and repeated design in healthy male subjects. The following treatments were to be administered under fasting conditions:

Treatment A (Test 1): One Fentanyl 1 mg/ml sublingual spray $(1\times100~\mu g~dose)$

27 Treatment B (Test 2): One Fentanyl 4 mg/ml sublingual spray $(1\times400 \mu g dose)$

Treatment C (Test 2): Two Fentanyl 4 mg/ml sublingual spray $(1\times800 \mu g dose)$

Treatment D (Placebo-Test 1): One Placebo 1 mg/ml sublin-5 gual spray (1×100 ug dose)

Treatment E (Placebo-Test 2): One Placebo 4 mg/ml sublingual spray (1×400 μg dose)

Treatment F (Placebo-Test 2): Two Placebo 4 mg/ml sublingual spray (1×800 μg dose)

The products were to be administered to nine (9) healthy male volunteers according to the following design in Table 16 below:

TABLE 16

	Period 1	Period 2	Period 3
Sequence 1 $(n = 6)$	100 μg	400 μg	800 µg
Sequence 2 $(n = 3)$	Placebo	Placebo	Placebo

In each period, subjects were to arrive at the clinical site at least 10 hours before dosing. After a supervised overnight fast, a single oral dose of the assigned formulation was to be orally administered in the morning. Subjects were allowed to 25 leave the clinical site after the 24-hour blood draw. The washout period was to be of at least 14 days; the duration of this study was expected to be approximately 5-6 weeks. As per

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protocol, each sequential dose was to be separated by a washout of at least 14 days, which corresponds to more than 10 time the expected half life of the moiety to be measured. However, during the study periods 1 and 2 were separated by a wash-out of 15 days and periods of 2 and 3 were separated by a wash-out of 13 days. As fentanyl's expected half life is reported to be approximately 6.4 hours, it is judged that these wash-out deviations should not affect the conclusion of the study. Furthermore, no carry-over was observed at the beginning of the second and third periods.

Pharmacokinetic Assessments

Blood samples for pharmacokinetic measurements were collected prior to and up to 24 hours (serial samplings) after each drug administration. The drug concentrations produced 15 by the administration of the studied formulations were used to derive the pharmacokinetic parameters listed in Tables 17 and

Six (6) subjects were included in the statistical analysis. A summary of the non-normalized pharmacokinetic parameters is presented in Table 17 and a summary of the normalized pharmacokinetic parameters is presented in Table 18. The mean measured plasma concentrations versus time profile, produced by the administration of the Test products, is depicted in FIG. 11, whereas the In-transformed mean concentrations versus time profile is presented in FIG. 12.

The pharmacokinetic parameters of interest for this study were C_{max} , AUC_{∞} , AUC_{T} , $AUC_{T/\infty}$, K_{el} , T_{max} , $T_{1/2el}$, Cl/F and V_Z/F .

TABLE 17

	Pharmacokinetic Parameters Fentanyl (n = 6) Non-normalized Data										
	TEST 1				TEST 2 (3 N =						
PARAMETER	MEAN	C.V. (%)	MEAN	C.V. (%)	MEAN	C.V. (%)					
C _{max} (pg/ml)	172.0	27.1	708.0	50.2	1270.4	37.7					
$In (C_{max}) (pg/ml)$	5.1207	4.8	6.4509	8.2	7.1102	5.4					
T _{max} (hours)	0.50	29.7	0.50	61.3	0.75	0.0					
$AUC_T(pg \cdot h/ml)$	472.6	66.2	3556.1	63.0	5417.3	30.6					
$\operatorname{In}\left(\operatorname{AUC}_{T}\right)\left(\operatorname{pg}\cdot\operatorname{h/ml}\right)$	6.0271	8.6	8.0208	7.5	8.5734	3.6					
AUC_{∞} (pg · h/ml)	817.9	36.1	4242.6	57.6	5726.8	28.8					
$\ln (AUC_{\infty}) (pg \cdot h/ml)$	6.6607	4.8	8.2303	6.4	8.6317	3.4					
$AUC_{T/\infty}$ (%)	54.90	28.7	81.48	10.6	94.35	1.8					
K_{el} (hour ⁻¹)	0.2008	27.4	0.1593	44.9	0.1782	0.9					
T _{1/2 el} (hours)	3.70	30.4	5.20	45.8	3.89	0.9					
Cl/F (ml/h/kg)	1718.8	27.7	1532.2	49.7	1837.3	13.1					
V _Z /F (ml/kg)	9070.2	34.7	10470.4	47.1	10307.7	12.2					

For T_{max} , the median is presented and the statistical analysis is based on ranks.

TABLE 18

	Da	ta Normaliz	ed to the 100 µg	g dose			
	TEST 1 (100 μg) n = 6		TEST 2 (- n =		TEST 2 (800 μg) N = 2		
PARAMETER	MEAN	C.V. (%)	MEAN	C.V. (%)	MEAN	C.V. (%)	
C _{max} (pg/ml)	172.0	27.1	177.0	50.2	158.8	37.7	
$\operatorname{In}\left(C_{max}\right)\left(\operatorname{pg/ml}\right)$	5.1207	4.8	5.0646	10.4	5.0307	7.7	
T _{max} (hours)	0.50	29.7	0.50	61.3	0.75	0.0	
$AUC_T(pg \cdot h/ml)$	472.6	66.2	889.0^{1}	63.0	677.2	30.6	
$\ln (AUC_T) (pg \cdot h/ml)$	6.0271	8.6	6.6346^{1}	9.1	6.4940	4.8	
AUC _m (pg·h/ml)	817.9	36.1	1060.7	57.6	715.9	28.8	
In (AUC _m) (pg · h/ml)	6.6607	4.8	6.8440	7.7	6.5523	4.5	
AUC _{7/∞} (%)	54.90	28.7	81.48^{2}	10.6	94.35^{2}	1.8	
K _{el} (hour ⁻¹)	0.2008	27.4	0.1593	44.9	0.1782	0.9	
T _{1/2 el} (hours)	3.70	30.4	5.20	45.8	3.89	0.9	

TABLE 18-continued

Data Normalized to the 100 μg dose									
		TEST 1 (100 µg) TEST 2 (400 µg) n = 6 n = 6			TEST 2 (800 μg) N = 2				
PARAMETER	MEAN	C.V. (%)	MEAN	C.V. (%)	MEAN	C.V. (%)			
Cl/F (ml/h/kg) V _Z /F (ml/kg)	1718.8 9070.2	27.7 34.7	1532.2 10470.4	49.7 47.1	1837.3 10307.7	13.1 12.2			

For T_{max} , the median is presented and the statistical analysis is based on ranks

¹= Different than Test - 1 (p < 0.05)

²= Different than Test - 1 (p < 0.01)

N.S. = Not Significant (p > 0.05)

Definition of the Pharmacokinetic Parameters

C_{max}: Maximum plasma concentration (ng/ml).

 T_{max} : Time of maximum measured plasma concentration; if it occurs at more than one time point, T_{max} is defined as the first time point with this value (hour).

 C_{LOC} : Last quantifiable concentration (ng/ml).

 T_{LQC} : Time of last quantifiable plasma concentration (hour). \overrightarrow{AUC}_T : Cumulative Area Under the plasma concentration time Curve calculated from 0 to T_{LQC} using the Trapezoidal method (ng·h/ml), and calculated according the following equation:

$$AUC_T = \sum_{2}^{N_{TLQC}} \left[\left(\frac{C_{n-1} + C_n}{2} \right) \times (T_n - T_{n-1}) \right]$$

Where N_{TLQC} is the number of the sample related to the T_{LQC} . AUC_{∞} : Extrapolated Area Under the plasma concentration time Curve to infinity (ng·h/ml).

$$AUC_{\infty} = AUC_T + \left(\frac{C_{LQC}}{K_{el}}\right)$$

 $\mathrm{AUC}_{T/\infty}$: Relative percentage of AUC_T with respect to AUC_∞ (%).

$$AUC_{T/\infty} = \left(\frac{AUC_T}{AUC_{\infty}}\right) \times 100$$

 T_{LIN} . Time point where log-linear elimination begins (hour). K_{el} : Apparent mean elimination rate constant estimated by a non-linear least-squares regression analysis; a minimum of three values are required at the end of the concentration-time curve (hour⁻¹).

 $T_{1/2el}$: Half-life of elimination (hour).

$$T_{1/2el} = \frac{\ln(2)}{K_{el}}$$

Cl/F: Apparent clearance (ml/h/kg)

$$Cl/F = \frac{Dose^*}{AUC_{\infty}}$$

 15 V_Z /F: Apparent volume of distribution (ml/kg)

$$V_Z/F = \frac{Dose^*}{K_{el} \cdot AUC_{\infty}}$$

*Dose expressed per weight (kg)

As noted in Table 11, the parameters for which a statistically significant difference was observed between the 100 μg dose and the normalized 400 μg dose was AUC_T , $\ln(AUC_T)$ and $AUC_{T/\infty}$ and the parameter for which a statistically significant difference was observed between the 100 μg dose and the normalized 800 μg dose was $AUC_{T/\infty}$. No statistically significant differences were observed for any of the other pharmacokinetic parameters under study.

The mean C_{max} value of the 100 μg dose was 172.0 pg/ml while for the 400 μg dose, it was 708.0 pg/ml and for the 800 μg dose, it was 1270.4 pg/ml. Once normalized to the 100 μg dose, the mean C_{max} value of the 400 μg dose was 177.0 pg/ml and for the 800 μg dose, it was 158.8 pg/ml.

The median T_{max} was 0.50 hour for the 100 µg dose and the 400 µg dose, and 0.75 hour for the 800 µg dose.

The mean AUC $_T$ value of the 100 $\mu \rm g$ dose was 472.6 pg·h/ml while for the 400 $\mu \rm g$ dose, it was 3556.1 pg·h/ml and for the 800 $\mu \rm g$ dose, it was 5417.3 pg·h/ml. Once normalized to the 100 $\mu \rm g$ dose, the mean AUC $_T$ value of the 400 $\mu \rm g$ dose was 889.0 pg·h/ml and for the 800 $\mu \rm g$ dose, it was 677.2 pg·h/ml.

The mean AUC_{∞} value of the 100 μg dose was 817.9 pg·h/ml while for the 400 μg dose, it was 4242.6 pg·h/ml and for the 800 μg dose, it was 5726.8 pg·h/ml. Once normalized to the 100 μg dose, the mean AUC_{∞} value of the 400 μg dose was 1060.7 pg·h/ml and for the 800 μg dose, it was 715.9 pg·h/ml.

The AUC_{T/∞} ratio was approximately 55% for the 100 pg dose, 81% for the 400 µg dose and 94% for the 800 µg dose.

The mean K_{el} was 0.2008 hour⁻¹ for the 100 μg dose, 0.1593 hour⁻¹ for the 400 μg dose and 0.1782 hour⁻¹ for the 800 μg dose, while the mean $T_{1/2el}$ value was 3.70 hours, 5.20 hours and 3.89 hours following the same order.

The mean Cl/F was 1718.8 ml/h/kg for the 100 μg dose, 1532.2 ml/h/kg for the 400 μg dose and 1837.3 ml/h/kg for the 800 μg dose, while the mean V_Z F was 9070.2 ml/kg for the 100 μg dose, was 10470.4 ml/kg for the 400 μg dose and was 10307.7 ml/kg for the 800 μg dose.

The intra-subject variation was 36.79%, 37.05% and 60 28.88% for C_{max} , AUC_T , and AUC_{∞} , respectively. Safety Evaluation

The safety parameters included the occurrence of adverse effects, measurements of vital signs, respiratory rate, oxygen saturation of blood by finger pulse oximetry, ECG and clini-65 cal laboratory parameters.

All adverse events were spontaneously reported by the volunteer, observed by the Clinical Investigator (or delegates)

or elicited by general questioning by the clinical staff. Adverse events were also reported upon completion of the form "Taste of Medication Questionnaire to Subject", which was filled right after dosing. For the purposes of the study, the period of observation for each individual subject extended from the time the subject gave informed consent to within 7 days following the last drug administration. Safety Results:

All nine subjects experienced a total of one-hundredtwenty-seven (127) adverse events during the study. No serious adverse events were recorded in this study. Twenty adverse events (8 different types) were reported after the single dose administration of the Test 1 (A) product, fifty-six adverse events (26 different types) were reported after the 15 single dose administration of the Test 2 (B) product, twentytwo adverse events (19 different types) were reported after the single dose administration of the Test 2 (C) product, ten adverse events (8 different types) were reported after the single dose administration of the Placebo-Test 1 (D) product, $_{20}$ eleven adverse events (9 different types) were reported after the single dose administration of the Placebo-Test 2 (E) product and twelve adverse events (8 different types) were reported after the single dose administration of the Placebo-Test 2 (F) product. Two (2) adverse events associated with 25 post-study laboratory test results were imputed to three formulations.

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Six subjects (100%) reported adverse events after the administration of the Test 1 (A) formulation, six subjects (100%) reported adverse events after the administration of the Test 2 (B) formulation, two subjects (100%) reported adverse events after the administration of the Test 2 (C) formulation, three subjects (100%) reported adverse events after the administration of the Placebo-Test 1 (D) formulation, three subjects (100%) reported adverse events after the administration of the Placebo-Test 2 (E) formulation and three subjects (100%) reported adverse events after the administration of the Placebo-Test 2 (F) formulation. The adverse events by system Organ Class are listed in Table 19.

The events abdominal distension, abdominal pain, abdominal pain upper, anxiety, depressed mood, diarrhea, disturbance in attention, dizziness (10 episodes out of 11), dry mouth, dry skin, dysgeusia, headache, fatigue (6 episodes out of 7), feeling cold, feeling drunk, feeling hot, feeling of relaxation, hot flush, hyperhidrosis, hypoaesthesia oral, hypoaesthesia, nasal congestion, nausea, oral discomfort, pallor, paresthesia oral, pruritus, sensation of heaviness, somnolence, speech disorder, tongue coated and vomiting were assessed to be possibly related to the drugs. The other events cough, dizziness (1 episode out of 11), fatigue (1 episode out of 7), musculoskeletal pain, rhinorrhoea and throat irritation were assessed to be not related to the study drugs. The other event nasopharyngitis was assessed to be unlikely related to the study drugs.

TABLE 19

		Sum	mary of A	dverse Ev	ents by Sy	stem Orga	ın Class					
		Peri	od 1			Peri	od 2			Peri	od 3	
Parameter		anyl) μg	Plac	cebo		Fentanyl 400 µg Placel		cebo	Fentanyl 800 µg		Plac	cebo
Number of subjects exposed		6		3 6			3	2		3		
Number of subjects reporting at least one adverse event		6		3		6		3		2		3
Total number of withdrawals		0		0		4		0		0		0
Withdrawals due to adverse event		0		0		1		0		0		0
(not related to tested drug) Withdrawals due to adverse event		0		0		2		0		0		0
(related to tested drug) Total number of adverse events*	2	0	1	.0	-	6	1	1	2	2	1	2
Adverse events at least possibly drug related*	_	8		8		3	_	1		2		2
Nature of adverse events	Number of subjects	% of subjects exposed										
			C	astrointes	tinal disor	ders						
Abdominal distension	0	0	0	0	1	17	0	0	0	0	0	0
Abdominal pain	0	0	0	0	0	0	1	33	0	0	1	33
Abdominal pain upper	0	0	0	0	1	17	0	0	0	0	0	0
Diarrhoea	0	0	0	0	0	0	1	33	0	0	1	33
Dry mouth	0	0	0	0	0	0	0	0	1	50	0	0
Dysgeusia	6	100	3	100	6	100	3	100	2	100	3	100
Hypoaesthesia oral	4	67	1	33	3	50	1	33	1	50	0	0
Nausea	0	0	0	0	4	67	0	0	1	50	0	0
Oral discomfort	1	17	0	0	1	17	1	33	1	50	0	0
Tongue coated	0	0	0	0	1	17	0	0	0	0	0	0
Vomiting	0	0 Gen	0 eral disord	0 lers and ac	1 Iministrati	17 on site coa	0 nditions	0	1	50	0	0
Fatigue	3	50	1	33	2	33	0	0	1	50	0	0
Feeling cold	0	0	0	0	1	17	0	0	1	50	0	0
Feeling drunk	0	0	0	0	6	100	0	0	0	0	0	0
Feeling hot	0	0	0	0	1	17	0	0	1	50	0	0
Feeling of relaxation	3	50	0	0	1	17	0	0	0	0	0	0
Hyperhidrosis	0	0	0	0	0	0	0	0	1	50	0	0

TABLE 19-continued

_		Peri	od 1			Perio	od 2			Perio	od 3	
Parameter	Fent 100	anyl) µg	Pla	cebo		tanyl) μg	Pla	cebo		tanyl 0 μg	Pla	cebo
				Investi	igations*							
Alanine aminotransferase increased Aspartate aminotransferase increased	0 0	0 0	1 1	33 33	0 0	0	1 1	33 33	0 0	0 0	1	33 33
		Mı	ısculoske	letal and co	onnective	tissue diso	rders					
Sensation of heaviness Musculoskeletal pain	0	0	0 0	0 0 Nervous sys	1 1 stem diso	17 17 rders	0 0	0 0	0 0	0	0	0
				•						400		
Dizziness Disturbance in attention	0	0	0 0	0	6 2	100 33	0	0	2 1	100 50	0	0
Headache Hypoaesthesia	0	0	0	0	1 4	17 67	0	0 0	0 1	0 50	1 0	33 0
Paresthesia oral Somnolence	0	0	1 1	33 33	3 1	50 17	1 0	33 0	0 2	0 100	3 1	100 33
Speech disorder	0	0	0	0 Psychiatr	0 ric disorde	0 ers	0	0	1	50	0	0
Anxiety	0	0	0	0	0	0	1	33	0	0	0	0
Depressed mood	0	0 Re	0 espiratory	0 , thoracic a	1 ind media	17 stinal disor	0 ders	0	0	0	0	0
Cough	1	17	0	0	0	0	0	0	0	0	0	0
Nasal congestion Nasopharyngitis	1 0	17 0	0 1	0 33	1 0	17 0	0 0	0 0	0	0 0	0	0
Rhinorrhoea Throat irritation	1 0	17 0	0 0	0 0	0 1	0 17	0 0	0 0	0	0 0	0 0	0
			Skin an	d subcutan	eous tissu	e disorders	S					
Dry skin	0	0	0	0	0	0	0	0	1	50	0	0
Pallor Pruritus	0 0	0 0	0 0	0 0	1 0	17 0	0	0 0	1 1	50 50	0	0 0
				Vascula	r disorder	s						
Hot flush	0	0	0	0	3	50	0	0	1	50	0	0

^{*=} Two (2) adverse events were associated with post-study laboratory test results and have been assigned to each period.

Conclusions

The pharmacokinetic parameters were well defined for the three doses ($100~\mu g$, $400~\mu g$ and $800~\mu g$) administered in this study. C_{max} and AUC_{∞} seem to be proportional, AUC_{T} is consistent with dose-proportionality between the $400~\mu g$ and $800~\mu g$. Furthermore, the two formulations of fentanyl (1 mg/ml and 4 mg/ml sublingual spray in doses of $100~\mu g$, $400~\mu g$ and $800~\mu g$) administered during the study were well tolerated in most of the subjects. No subject participating in the trial reported serious adverse events during the course of this study.

EXAMPLES 8-12

In-vitro Permeation Experiments

The permeation characteristics of fentanyl formulations were studied using EpiOral tissues (ORL-100). MatTek Corp's EpiOral is used as model for screening sublingual 60 drug absorption of pharmaceutical formulations. MatTek's EpiOral tissue consists of normal, human-derived epithelial cells. The cells have been cultured to form multilayered, highly differentiated models of the human buccal (EpiOral) phenotypes. The EpiOral tissue model exhibits in vivo-like 65 morphological and growth characteristics which are uniform and highly reproducible. Morphologically, the tissue model

closely parallels native human tissue, thus providing a useful in-vitro means to assess in-vivo permeability of pharmaceutical formulations across sublingual mucosa.

The EpiOral tissues, grown on cell culture inserts with Teflon backing membrane, were shipped by MatTek Corp on Monday for delivery on Tuesday morning. All the tissues were used in the permeability experiments within 72 hours of shipment. The inserts containing the tissues were rinsed with distilled water before the start of permeation experiments. The tissue area for the ORL-100 is 0.6 cm².

The receiver compartment (wells) received 0.3 ml of phosphate citrate buffer of pH 6.6 (receiver solution). The donor compartments (tissue inserts) were placed in the wells and filled with 0.5 ml of drug solution (donor solution).

The inserts were moved from well to well containing fresh receiver fluid at pre-determined intervals (2, 5, 7, 9, 11, 13, 15, 30, 45, 60, 90, and 120 min). After permeation studies both donor and receiver fluids were collected in vials for analysis by HPLC. The receiver and donor solution concentrations and the flux over each permeation time interval were determined.

EXAMPLE 8

In Example 8, the permeation of fentanyl base was compared to the permeation of fentanyl citrate. The formulations and percent permeated in 2 hours is listed in the table below:

TABLE 20

	EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	MIGLYOL % (V)	% PERMEATED IN 2 HOURS
Fentanyl Base	8-a (b)	1 mg/ml	20	5	_	17.33
Fentanyl Base	8-b (w)	1 mg/ml	20	5	_	17.18
Fentanyl Citrate	8-c (b)	0.646 mg/ml	20	5	_	1.81
Fentanyl Base	8-d	1 mg/ml	20	5	79.3	1.64

(b)—buffer,

(w)—water

EXAMPLE 9

In Example 9, Effect of Alcohol on the Permeation of Fentanyl Base Formulations was tested. The formulations and percent permeated in 2 hours are listed in the Table below:

TABLE 21

EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	% PERMEATED IN 2 HOURS
9-a (w) (Control)	1 mg/ml	20	5	17.18
9-b (w)	1 mg/ml	28	5	13.45
9-c (w)	1 mg/ml	40	5	10.95
9-c (w)	1 mg/ml	50	5	9.18

(b)—buffer, (w)—water

EXAMPLE 10

In Example 10, the effects of PG (propylene glycol) on Fentanyl Formulations was tested. The formulations and percent permeated in 2 hours are listed in the table below:

TABLE 22

EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	% PERMEATED IN 2 HOURS
10-a (w) (Control)	1 mg/ml	20	5	17.18
10-b (w)	1 mg/ml	20	25	14.518

(b)—buffer,

EXAMPLE 11

In Example 11, the effect of pH on Fentanyl Formulations was tested. The formulations and percent permeated in 2 hours are listed in the table below. As indicated in the table, the permeation of fentanyl across buccal tissue was dependent on the pH of the formulation. Because fentanyl has pKa value (7.3 and 8.4) within the pH range studied, its degree of ionization changed as the pH of the formulation was altered. The results in the table below indicate that that formulations adjusted to pH between 8 and 9 were showing better permeability and physical stability.

TABLE 23

30	EXAM- PLE #*	CONC. OF FENTANYL BASE	BUFFER PH	ALCOHOL % (V)	PG % (V)	% PERMEATED IN 2 HOURS
	11-a (w) (Control)	2 mg/ml	Water	30	5	17.83
35	11-b (b)	2 mg/ml	5.5	30	4	5.33
	11-c (b)	2 mg/ml	6.6	29	4	10.62
	11-d (b)	2 mg/ml	8.6	29	4	13.48
	11-e (b)	2 mg/ml	9.6	29	4	11.8

(b)—buffer, 40 (w)—water

EXAMPLE 12

In Example 12, several ingredients including hydroxypropyl beta cyclodextrin (HPBCD), mannitol, polyvinyl pyrrolidone (PVP), propylene carbonate (PC), sodium glycocholate
(SG), sodium lauryl sulphate (SLS), triacetin, triethyl citrate
and tween 80 (TW 80) were added to the formulations either
individually or in combination and studied for their effect on
permeability and solution stability. Table 24 to 36 summarizes the formulations and permeation results of buffered and
water formulations containing the above excipients.

TABLE 24

	Results of the effects of Buffer and Water on Fentanyl Formulations.											
CONC. OF					INACTIV	-						
EXAMPLE #*	FENTANYL BASE	BUFFER OR WATER	ALCOHOL % (V)	PG % (V)	Mannitol % (wt)	Triacetin % (wt)	TW80 % (wt)	% PERMEATED IN 2 HOUR				
12-a (b) (Control)	1 mg/ml	В	20	5	_	_	_	17.33				
12-b (w) (Control)	1 mg/ml	W	20	5	_	_	_	17.18				
12-c (b)	1 mg/ml	В	20	5	0.1	0.15	_	16.14				
12-d (w)	1 mg/ml	W	20	5	0.1	0.15	_	12.10				
12-e (b)	1 mg/ml	В	20	5	0.3	_	0.2	16.02				

TABLE 24-continued

	Results of the effects of Buffer and Water on Fentanyl Formulations.								
	CONC. OF					/E INGRED	-		
EXAMPLE #*	FENTANYL BASE	BUFFER OR WATER	ALCOHOL % (V)	PG % (V)	Mannitol % (wt)	Triacetin % (wt)	TW80 % (wt)	% PERMEATED IN 2 HOUR	
12-f (w)	1 mg/ml	W	20	5	0.3	_	0.2	12.35	
12-g (b)	1 mg/ml	В	20	5	0.15	0.3	0.1	16.43	
12-h (w)	1 mg/ml	W	20	5	0.15	0.3	0.1	9.76	
12-i (b)	1 mg/ml	В	20	5	0.2	_	0.2	18.74	
12-j (w)		W	20	5	0.2	_	0.2	13.19	
12-k (b)	1 mg/ml	В	20	5	_	_	0.3	14.73	
12-l (w)	1 mg/ml	W	20	5	_	_	0.3	13.76	

(b)—buffer,

(w)—water

The results in Table 24 indicate that all the buffered formulations had similar permeability characteristics as that of control formulations except the buffered formulation containing

0.3% Tween 80 which showed lower permeability. All water formulations exhibited lower permeability than buffered formulations.

TABLE 25

Effect of HPBCD on Fentanyl Formulations							
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	HPBCD (MOLECULAR RATIO TO API)	% PERMEATED IN 2 HOURS		
12-m (w) (Control)	1 mg/ml	20	5	_	17.18		
12-n (w) 12-o (w)	1 mg/ml 1 mg/ml	20 20	5 5	1:1 (0.413%) 1:2 (0.826%)	14.28 13.50		

(b)—buffer,

(w)—water

TABLE 26

	Effect of Mannitol on Fentanyl Formulations								
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	MANNITOL % (WT)	% PERMEATED IN 2 HOURS				
12-p (b) (Control)	1 mg/ml	20	5	_	17.33				
12-q (b)	1 mg/ml	20	5	0.3	17.71				
12-r (b)	1 mg/ml	20	5	0.4	16.86				
12-s (b)	1 mg/ml	20	5	0.5	15.41				
12-t (b)	1 mg/ml	20	5	0.8	14.81				

(b)—buffer,

(w)—water

TABLE 27

EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	entanyl Formulat PVP % (WT)	% PERMEATED IN 2 HOURS
12-u (w) (Control)	1 mg/ml	20	5	_	17.18
12-v (w)	1 mg/ml	20	5	0.1	16.73
12-w (w)	1 mg/ml	20	5	0.5	14.68
12-x (w)	1 mg/ml	20	5	1	14.52
12-y (w)	1 mg/ml	25	5	3	10.75

(b)—buffer,

(w)-water

TABLE 28

EXAMPLE#*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	PC % (WT)	% PERMEATED IN 2 HOURS
12-z (w) (Control)	1 mg/ml	20	5	_	17.18
12-aa (w)	1 mg/ml	20	5	1	14.39
12-bb (w)	1 mg/ml	20	5	1.5	14.43

(b)—buffer,

(w)—water

TABLE 29

E	Effect of Sodium Glycocholate (SG) on Fentanyl Formulations							
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	SG % (WT)	% PERMEATED IN 2 HOURS			
12-cc (w) (Control)	1 mg/ml	20	5	_	17.18			
12-dd (w) 12-ee (w)	1 mg/ml 1 mg/ml	20 20	5 5	0.5 1	18.30 19.78			

(b)—buffer, (w)—water

TABLE 30

	Effect of Triacetin on Fentanyl Formulations							
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	TRIACETIN % (WT)	% PERMEATED IN 2 HOURS			
12-ff (b) (Control)	1 mg/ml	20	5	_	17.33			
12-gg (b) 12-hh (b)	1 mg/ml 1 mg/ml	20 20	5 5	0.5 2	17.71 14.56			

(b)—buffer, (w)—water

TABLE 31

	Effect of Triethyl Citrate on Fentanyl Formulations							
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	TRIETHYL CITRATE % (WT)	% PERMEATED IN 2 HOURS			
12-ii (w) (Control)	1 mg/ml	20	5	_	17.18			
12-jj (b) 12-kk (b)	1 mg/ml 1 mg/ml	20 20	5 5	0.5 2	15.81 10.02			

(b)—buffer, (w)—water

TABLE 32

Effect of Tween 80 (TW 80) on Fentanyl Formulations							
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	TW80 % (WT)	% PERMEATED IN 2 HOURS		
12-ll (b) (Control)	1 mg/ml	20	5	_	17.33		

TABLE 32-continued

Effect of Tween 80 (TW 80) on Fentanyl Formulations							
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	TW80 % (WT)	% PERMEATED IN 2 HOURS		
12-mm (b) 12-nn (b)	1 mg/ml 1 mg/ml	20 20	5 5	0.3 0.6	14.73 13.72		

(b)—buffer, (w)—water

TABLE 33

Effect of Labrasol & SLS on Fentanyl Formulations								
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	LABRASOL % (WT)	SLS % (WT)	% PERMEATED IN 2 HOURS		
12-oo (w) (Control)	1 mg/ml	20	5	_	_	17.18		
12-pp (w)	1 mg/ml	20	5	6.5	_	8.25		
12-qq (w)	1 mg/ml	20	5	3.0	0.5	9.31		
12-rr (w)	1 mg/ml	20	5	6.5	0.5	8.63		

(b)—buffer, (w)—water

 $TABLE\ 34$

	Effect of Mannitol & Triacetin on Fentanyl Formulations								
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	MANNITOL % (WT)	TRIACETIN % (WT)	% PERMEATED IN 2 HOURS			
12-ss (b)	1 mg/ml	20	5	_	_	17.33			
(Control)									
12-tt (b)	1 mg/ml	20	5	0.1	0.15	16.14			
12-uu (b)	1 mg/ml	20	5	0.15	0.4	18.33			
12-vv (b)	1 mg/ml	20	5	0.15	0.5	17.14			
12-ww (b)	1 mg/ml	20	5	0.2	0.3	16.73			
12-xx (b)	1 mg/ml	20	5	0.25	0.5	16.31			
12-yy (b)	1 mg/ml	20	5	0.3	0.2	17.70			
12-zz (b)	1 mg/ml	20	5	0.4	0.2	16.88			

(b)—buffer, (w)—water

TABLE 35

Effect of Mannitol & TW 80 on Fentanyl Formulations							
CONC. OF							
12-aaa (b) (Control)	1 mg/ml	20	5	_	_	17.33	
12-bbb (b)	1 mg/ml	20	5	0.2	0.2	18.74	
12-ccc (b)	1 mg/ml	20	5	0.3	0.1	16.53	
12-ddd (b)	1 mg/ml	20	5	0.3	0.2	16.02	

(b)—buffer, (w)—water

TABLE 36

Effect of Triacetin & TW 80 on Fentanyl Formulations							
EXAMPLE #*	CONC. OF FENTANYL BASE	ALCOHOL % (V)	PG % (V)	TW 80 % (WT)	TRIACETIN % (WT)	% PERMEATED IN 2 HOURS	
12-eee (b) (Control)	1 mg/ml	20	5	_	_	17.33	
12-fff (b) 12-ggg (b)	1 mg/ml 1 mg/ml	20 20	5 5	0.1 0.1	0.2 0.3	16.99 16.63	

(b)—buffer,

The results indicate that addition of individual excipients 15 (with 1 mg/ml of fentanyl base) containing less than 20% of including HPBCD (Table 25), PVP (Table 27), PC (Table 28), Triethyl citrate (Table 31) and Tween 80 (Table 32) to the formulation decreased the permeability of fentanyl across MatTek buccal membranes irrespective of excipient concentration. As shown in Table 26 and Table 30, formulations 20 containing 0.3% Mannitol and 0.5% Triacetin showed similar permeability characterisities as that of control formulation but the permeability decreased as the concentrations of these individual excipients were increased in the formulations. Stability studies indicated that a minimum of 0.45% and 0.5% 25 mannitol concentration should be added to buffer and water formulations, respectively, to keep them stable. In case of Triacetin, formulations containing 0.5% or higher concentrations of triacetin were found to be stable.

Permeation of fentanyl from formulations containing SG 30 was greater than that of control formulations (Table 29). Also, we observed that fentanyl permeation increased with the increase of SG concentration in the formulation.

The results of fentanyl permeation across MatTek-buccal tissues from formulations containing combination of excipi- 35 ents are shown in Tables 33-36. Addition of Labrasol to the formulation improved the stability but decreased fentanyl permeation across MatTek buccal tissues. Similar results were observed with formulations containing labrasol and SLS. Among all the formulations containing combination of 40 excipients, two formulations, Example 12-uu (0.15% mannitol, 0.4% triacetin) and Example 12-bbb (0.2% mannitol, 0.2% TW80), showed higher permeability compared to control formulation. We observed that the presence of mannitol in triacetin formulations did not show any improvement in the 45 permeation. Hence, the formulation containing 0.5% triacetin (Table 30) was selected for further studies. Though the formulation, Example 12-bbb, showed good permeability mannitol concentration was increased to 0.3% to improve the stability of the product.

Both fentanyl citrate and fentanyl base formulations were stable at all temperatures studied. The data from in-vitro tissue permeation studies, as shown in Table 3, showed that permeation of fentanyl from fentanyl base formulations was about 10-fold higher than from fentanyl citrate formulations. 55 Water and buffer formulations did not show any significant difference in fentanyl base permeation across buccal tissue. Our studies also showed that fentanyl base formulation containing Miglyol had very low permeability. Among the excipients, triacetin at 0.5% and mannitol at 0.3% in combination with 0.2% TW 80 showed good permeability and stability.

The transport of fentanyl across buccal tissues was dependent on ethyl alcohol (ethanol) and propylene glycol (PG) concentrations in the formulation. Ethanol is used as a cosol- 65 vent to solubilize fentanyl base in aqueous based formulations. In this investigation, we observed that formulations

ethanol were precipitating at refrigerated conditions. Also, we observed that permeation of fentanyl across buccal tissue was indirectly proportional to ethanol concentrations in the formulations (Table 21). Similar results were observed with PG (Table 22). PG was helpful in increasing the solubility of fentanyl base in aqueous solution and also in enhancing the permeation of the fentanyl base across buccal tissues. Formulations containing more than 5% PG did not show significant improvement in the permeation of fentanyl, however, the solution viscosities increased proportionally with PG concentrations. The formulations containing more than 5% PG did not show good spray characteristics.

EXAMPLE 13

In Example 13, a formulation having the following ingredients in Table 37 was tested. The formulation did not include fentanyl.

TABLE 37

Ingredient	
Dehydrated Alcohol % (v/v)	30%
Propylene glycol % (v/v)	5%
Peppermint Oil % (v/v)	0.2%
Borate buffer (pH 9)	QS
Hydroxy Propyl Beta Cyclodextrin (HPBCD)	1:2

The formulation was sprayed using a 0.10 ml multidose nasal spray pump by Pfeiffer of America, Princeton, N.J. and the droplets were measured using a Malvern Mastersizer S device, by Malvern Instruments Ltd. A single depression of the sublingual spray pump generated a plume which was then analyzed for spray particles. The sample size for the dose volume, spray pattern, and droplet size distribution was 25 sprays.

Droplet Volume

In the droplet volume evaluation, 25 spray samples were evaluated using 5 different stroke numbers for each spray sample the following results were measured:

Overall mean=100.4 µl

Maximum single actual value=103.2 μl

Lowest single actual value=95.3 µl

Standard deviation=1.1

Range=7.9

Coefficient of variation=1.1%

Spray Pattern

In the spray pattern evaluation, 25 spray samples were evaluated using a manual actuation at 30 mm from the target. The formulation was dyed with methylene blue and the following spray pattern results were measured:

Small Diameter [mm]

min: 35.4

46

mean: 50.6
max: 62
s: 7.00
Largest Diameter [mm]
min: 40
mean: 56.9
max: 67
s: 6.01
Spray Angle
min: 64°
mean: 83.3°
max: 94°
s: 7.03
Ratio (Largest/Smallest Diameter)
min: 1.04
mean: 1.13
max: 1.33
s: 0.073
Droplet Size Distribution
In the droplet size distribution evaluation, 25 spray samples
were evaluated using a manual actuation at 30 mm from the
target. The following droplet size distribution results were
measured:
Percentage Share of Droplet Diameters at 10 μm [%]
min: 0.65
mean: 1.66
max: 2.70
s: 0.527
10% of the Droplet Diameters are Smaller than the Indicated
Value [µm]
min: 15
mean: 18.2
max: 23
s: 1.91
50% of the Droplet Diameters are Smaller than the Indicated
Value [µm]
min: 35
mean: 44.7
max: 65
s: 7.52
90% of the Droplet Diameters are Smaller than the Indicated Value [µm]
min: 96
mean: 154.4
max: 349
s: 64.42
EXAMPLE 14

EXAMPLE 14

In Example 14, a formulation having the following ingredients in Table 38 was tested. The formulation did not include fentanyl.

TABLE 38

Ingredient	
Dehydrated Alcohol % (v/v)	30%
Propylene glycol % (v/v)	5%
Peppermint Oil % (v/v)	0.2%
Borate buffer (pH 9)	QS
Hydroxy Propyl Beta Cyclodextrin (HPBCD)	1:2
Hydroxy Propyl Cellulose (HPC EP) % (v/v)	1%

The formulation was sprayed using a 0.10 ml multidose nasal spray pump by Pfeiffer of America, Princeton, N.J. and the droplets were measured using a Malvern Mastersizer S device, by Malvern Instruments Ltd. A single depression of the sublingual spray pump generated a plume which was then analyzed for spray particles. The sample size for the dose volume, spray pattern, and droplet size distribution was 25 sprays.

Droplet Volume

In the droplet volume evaluation, 25 spray samples were evaluated using 5 different stroke numbers the following results were measured:

Overall mean value=101.5 µl

Maximum single actual value=103.7 μl

15 Lowest single actual value=96.1 μl

Standard deviation=1.3

Range=7.6

Coefficient of variation=1.2%

Spray Pattern

In the spray pattern evaluation, 25 spray samples were evaluated using a manual actuation at 30 mm from the target. The formulation was dyed with methylene blue and the following spray pattern results were measured:

Small Diameter [mm]

25 min: 29 mean: 34.0 max: 46 s: 3.47

Largest Diameter

30 min: 35 mean: 40.8 max: 55 s: 4.13 Spray Angle 35 min: 58° mean: 63.8° max: 80° s: 4.65

Ratio (Largest/Smallest Diameter)

40 min: 1.08 mean: 1.20 max: 1.50 s: 0.092

Droplet Size Distribution

In the droplet size distribution evaluation, 25 spray samples were evaluated using a manual actuation at 30 mm from the target. The following droplet size results were measured:

Percentage Share of Droplet Diameters at 10 µm [%]

min: 0.24 50 mean: 0.68 max: 1.28 s: 0.278

10% of the Droplet Diameters are Smaller than the Indicated Value $[\mu m]$

55 min: 22 mean: 26.7 max: 35 s: 2.95

60

50% of the Droplet Diameters are Smaller than the Indicated

Value [µm] min: 61 mean: 83.3 max: 114 s: 11.41

5 90% of the Droplet Diameters are Smaller than the Indicated Value [μm]

min: 192

mean: 294.6 max: 440 s: 53.32

EXAMPLE 15

In Example 15, an assay procedure for fentanyl in 1, 2, 4, 6, and 8 mg/mL fentanyl sublingual spray samples with a working concentration of between 0.1 μ g/mL and 5 μ g/mL fentanyl in solution was performed. This method was developed and qualified in compliance with GMP requirements. The method was determined to be linear over the range of 0.05 μ g/mL to 7.83 μ g/mL fentanyl. The fentanyl working standard solution was stable over a seven day period in volumetric glassware and amber HPLC vials at refrigerated and ambient 15 conditions.

The equipment and supplies utilized in this process included an HPLC system equipped with a pump, variable wavelength detector, and autosampler, or equivalent, a Waters Symmetry HPLC column (C18, 4.6×75 mm, 3.5 μ m particle 20 size), 0.45 μ m, 47 mm nylon filters (Gelman Nylaflo® P/N 66608 or equivalent), acetonitrile (HPLC Grade), potassium phosphate monobasic (ACS Grade), phosphoric acid (ACS Grade), deionized water, alcohol (ethanol, absolute), and fentanyl base reference standard.

The solution preparations were prepared as described below and may be scaled as required.

For each liter of phosphate buffer solution (50 mM $\rm KH_2PO_4$ pH 2.8), 6.8 g of potassium phosphate mono basic and 1 liter of water was combined in a suitable vessel and 30 mixed well. The pH of the solution was adjusted to 2.8 with the drop-wise addition of phosphoric acid. The solution was filtered through 0.45 μ m nylon. This solution expires after one month.

For each liter of mobile phase (25% ACN, 75% phosphate 35 buffer) solution, 750 mL of phosphate buffer solution was combined with 250 mL of acetonitrile in a suitable vessel and mixed well. The system is degassed by an appropriate method before use if required. This solution expires after one month.

For each liter of stock diluent (95/5, Ethanol/Acetonitrile) 40 solution, 950 mL ethanol and 50 mL acetonitrile was combined in a suitable container and mixed well. This solution expires after one month.

Stock standard I, 40 µg/mL, ("SSI"), was prepared by weighing approximately 10.0 mg of fentanyl reference standard and added to a 250 mL volumetric flask. Approximately 200 mL stock diluent was added and swirled to dissolve the solid material. The mixture was diluted to the desired volume with stock diluent and mixed well.

Stock standard II, 40 µg/mL, ("SSII"), was prepared by 50 weighing approximately 10.0 mg of fentanyl reference standard and add to a 250 mL volumetric flask. Approximately 200 mL of stock diluent was added and swirled to dissolve solid material. The mixture was diluted to the desired volume with stock diluent and mixed well.

Working standard I, 2.4 μ g/mL, ("WSI"), was prepared by transferring 3.0 mL of stock standard I to a 50 mL volumetric flask. The mixture was diluted to the desired volume with mobile phase solution and mixed well. This solution expires after seven days.

Working standard II, 2.4 μ g/mL, ("WSII") was prepared by transferring 3.0 mL of stock standard II to a 50 mL volumetric flask. The mixture was diluted to the desired volume with mobile phase solution and mixed well. This solution expires after seven days.

The chromatographic conditions for the HPLC procedure are set forth below:

Column: HPLC Column Waters Symmetry C18, 4.6 x 75 mm, 3.5 um particle size

UV Detection: 214 nm Flow Rate: 2.0 mL/minute Injection Volume: 50 µL

Injection Volume: 50 μ L Temperature: Ambient (The temperature may be controlled

at 25° C.) Acquisition Time: 13 minutes

HPLC Procedure

After the system suitability is established, a maximum of 12 sample solutions can be injected in between working standards. A typical sequence would be as follows:

2× Mobile Phase (Blank)

2× Working standard II

5× Working standard I

Ix Sample (up to 12 injections)

Ix Working standard I

Ix Sample (up to 12 injections)

I× Working standard I

System Suitability

There should be no significant interfering peaks present at the retention time of fentanyl in the mobile phase blank injections. In terms of injection precision, the RSD of fentanyl for five replicate injections of working standard I should not exceed 2.0%. In terms of standard agreement, the agreement between the average peak response for the first five working standard I injections and the two working standard II injections should be between 98 to 102%. The agreement between working standards I and II need only be demonstrated once during the expiry of standards. The tailing factor at 5% peak height for fentanyl in the first working standard I injection should be between 0.8 and 1.5. In terms of standard precision over the run, the RSD of peak area for fentanyl in the working standard I injections over the run (OTR) should not exceed 2.0%.

Calculations are performed as set forth below. Working standard concentration is calculated as follows:

Mass of 1000 standard (mg)
$$*\frac{\mu g}{1 \text{ mg}} *\frac{3.0 \text{ ml}}{50.0 \text{ ml}} * \text{ of } = \mu g/\text{mL}$$
 Fentanyl Standard

Response Factor (R_f) is calculated as follows:

Fentanyl Peak
$$\frac{\text{Area}}{\text{Fentanyl}} *= R_f$$
 Concentration
$$(\mu g / \text{mL})$$

Standard Agreement is calculated as follows:

Average
$$R_f$$
 WSII $*100 = \%$ Standard Agreement

EXAMPLE 16

In Example 16 the method for determination of droplet size distribution by laser diffraction for fentanyl sublingual spray using the Spraytec device by Malvern was performed.

All data generated and described within this report were reviewed for compliance with Good Manufacturing Practices (21 CFR Parts 210 and 211).

The purpose of this project was to develop and validate a droplet size distribution method by laser diffraction for use with fentanyl sublingual spray product and placebo. The first portion of the project performed product evaluations to determine the proper automated actuation parameters to be used with the MightyRunt Actuation Station by Innova Systems, Inc. Using the automated actuation station, development of the droplet size distribution method for the sublingual product included vignetting studies, exhaust placement studies, and device placement studies.

The method validation evaluated intermediate precision between two analysts performing the developed method. All method development and qualification activities were performed using placebo.

Samples were prepared using Pfeiffer unit dose glass vials, Pfeiffer unit dose VI stoppers, Pfeiffer vial holder, and Pfeiffer unit dose applicator. The instrumentation utilized in the study include a Spraytec with 200 mm lens by Malvern Instruments, Inc, a MightyRunt Actuation Station by Innova Systems Inc. equipped with an exhaust fan attachment, and a Mettler Toledo balance Model AT201.

Actuation Parameter Study

Using the Spraytec to track the plume duration and droplet size distribution, the actuation parameters for the MightyRunt Actuation Station (MightyRunt) were optimized to replicate the plume duration, droplet size distribution, and shot weight generated by manual actuation. Dv10 (10% of the droplet diameters are smaller that the indicated value), Dv50 (50% of the droplet diameters are smaller that the indicated value), and Dv90, (90% of the droplet diameters are smaller that the indicated value), results from six devices with manual actuations were compared with the results from the six devices with automated actuations.

Acceptance Criteria

The individual shot weight results for the automated actuations should all fall within the range of 75%-125% of the average shot weight for the manual actuations. The average Dv10, Dv50 and Dv90 results of the automated actuations 45 should be within 75% to 125% of the average Dv10, Dv50 and Dv90 results for the manual actuations.

Statistical analysis shall include performance of a students' t-test on the two sets of droplet size distribution results. The results of the students' t-test should indicate that the manual 50 versus automated sets of data is statistically equivalent.

Method Development

Method development involved a vignetting study and 55 exhaust study utilizing the Spraytec. Actuations were performed using the MightyRunt and previously determined parameters. Vignetting occurs during laser diffraction analysis when the small droplets of a spray plume scatter the laser at an angle too steep to be captured by the range lens. Placement of the device close enough to the range lens to capture all of the scattered light without deposition on the range lens is critical. The vignetting study determined the appropriate range lens for analysis and the appropriate distance from device to range lens.

Exhaust placement affects plume travel. Plume velocity should not be accelerated by the draw of the exhaust and large

50

droplets should not fall while traveling through the laser path. The exhaust study determined the appropriate position behind the plume of the spray to ensure proper plume capture after passing through the laser path. The method included two distances for analysis from the tip of the nozzle to the path of the laser for a more complete characterization of the droplet size distribution.

Method Validation

Validation consisted of determining the precision and ruggedness of the method. A total of 24 devices from a single lot of placebo were used in the validation. One analyst tested 6 actuations at each distance. To demonstrate ruggedness, a second analyst repeated the analyses. Dv10, Dv50, and Dv90 results were compared.

Acceptance Criteria

The individual shot weight results for analyst one should all fall within the range of 75%-125% of the average shot weight for analyst two. The average Dv10, Dv50 and Dv90 results for analyst one should be within 75% to 125% of the average Dv10, Dv50 and Dv90 results for analyst two.

The final experimental procedure is set forth below.

Prior to analysis, the background and scattering profiles were verified as appropriate for analysis.

The sample bottle is inserted into the nozzle holder. Coasters were used to raise the platform for minimal adjustment. The coaster attached to the device holder was placed on top of coasters used for adjustment.

The MightyRunt with bottle and exhaust fan was placed in the appropriate positions for analysis. The MightyRunt was raised with the nozzle centered in front of the laser path with a lab jack. It was ensured that the MightyRunt was level following adjustment. Two bottle-to-laser distances were be evaluated, 7 cm and 4 cm, measured from the pump tip to the center of the laser path. For all analyses, the bottle was placed 14 cm from the range lens support structure to the pump tip.

The exhaust fan was turned on and placed on a lab jack 3 cm behind the laser path, centered behind the device, measured from the center of the laser path to the front edge of the exhaust shield. The exhaust fan had an impaction surface for the droplets to adhere to, i.e. a cheesecloth placed in the path of the droplets. 1/8 sheet of 18×36 inch cheesecloth folded into approximately a 4.5-inch square (4 layers of cheesecloth) is a sufficient impaction surface and will not impede the flow of the fan.

The device was actuated using the MightyRunt station and droplet size distribution was collected for each individual shot at the appropriate distances for analysis.

A report was printed for the entire plume duration and the plume plateau.

Data Reporting

The data for the D10, D50, D90, and Span values for each actuation are reported. The average and precision (% RSD) (percent relative standard deviation) were calculated for the D10, D50, and D90. Note: Span is defined as (D90–D10)/D50. Individual droplet size distribution results (μ m) and span values (unitless) were reported to X.XX. All average droplet size distribution results should be reported to X.X μ m. All RSD values should be reported to X.X %.

Statistical analysis included performance of a students' t-test on the two sets of droplet size distribution results. The

results of the students' t-test should have indicated that the analysts sets of data is statistically equivalent.

Results and Discussion

All spray plumes have three stages. The first stage, development, was characterized by variable droplets and decreasing transmission of the laser. The second stage, stable, was characterized by a stable droplet distribution and transmission. Variable droplets and increasing transmission characterize the final stage, dissipation. All comparisons of droplet size distribution used the stable stage of the plume.

Actuation Parameter Study

During the optimization of the actuation parameters for the MightyRunt, the plumes from manual and automated actuations were measured using the Spraytec. Dv10, Dv50, and Dv90 results were compared to optimize the automated parameters. The type of device that was examined in this study was significantly different from typical nasal sprays. Rather than a spring requiring a consistent force to initiate actuation and deliver the drug, there was an amount of force that was required to break the tabs or actuate the device. This force was not the same amount of force required to deliver the drug from the device once the tabs are broken. Because this device did not contain a spring, the parameters of interest 30 were actuation force, force rise time, and minimum travel distance. The finalized MightyRunt parameters are listed below in Table 39 and the utilized Spraytec settings are listed below in Table 40.

TABLE 39

Mighty Runt Actuation Parameters			
Parameter	Setting		
Actuation Force	4.0 kg		
Force Rise Time	0.2 sec		
Hold Time	1.0 sec		
Force Fall Time	1.0 sec		
Spray Delay	1 sec		
Minimum Travel Distance	10 mm		
Maximum travel Time	4.0 sec		
Trigger Signal Delay	0.0 sec		
Stage	Yes		

TABLE 40

Spraytec Settings			
Option	Setting		
Test Duration	200 msec		
Data Acquisition Rate	1000 Hz		
Acquisition Duty Cycle	0%		
Experimental Trigger	Transmission		
Transmission Trigger	98%		
Range Lens	200 mm		

The comparison of manual and automated actuations was performed using six devices with manual actuations and six devices with automated actuations. The droplet size distribution and shot weight results are summarized in Table 41 below.

52 TABLE 41

5	Device	D10 (µm)	D50 (μm)	D90 (µm)	Shot Weight (mg)	% Average
5	Manual 1	19.0	44.5	76.8	89.1	
	Manual 2	18.1	42.7	84.9	86.9	
	Manual 3	33.3	58.7	108.4	90.4	
	Manual 4	20.8	44.6	81.2	87.0	
	Manual 5	18.3	42.7	80.7	87.8	
10	Manual 6	19.0	44.3	81.5	84.9	
	Average	21.4	46.3	85.6	87.7	
	% RSD	27.5	13.3	13.4	2.2	
	Mighty Runt 1	17.3	37.5	65.1	83.8	96
	Mighty Runt 2	16.1	35.3	64.0	66.0	75
	Mighty Runt 3	18.8	38.9	66.2	84.2	96
15	Mighty Runt 4	15.4	37.8	77.6	85.4	97
13	Mighty Runt 5	17.6	38.6	67.1	82.4	94
	Mighty Runt 6	19.4	41.9	74.8	88.0	100
	Average	17.4	38.3	69.1	81.6	
	% RSD	8.8	5.6	8.2	9.7	
	Overall	19.43	42.30	77.36	84.7	
	average					
20	Overall % RSD	23.7	14.3	15.7	7.5	
	Automated	81	83	81		
	Actuation					
	Average as %					
	of Manual					
	Actuation					
25	Average					
	t-stat	1.60	2.97	3.15		
	t-critical	1.94	1.94	1.81		
	Result	Same	Different	Different		
		population	population	population		

Acceptance Criteria

Shot weight results for the automated actuations ranged 35 from 75%-100% of the average for manual actuations, meeting the acceptance criteria of 75%-125% of the average shot weight for the manual actuations. The average Dv10 for automated actuations was 81% of the average Dv10 for manual actuations. The average Dv50 for automated actuations was 40 83% of the average Dv50 for manual actuations. The average Dv90 for automated actuations was 81% of the average Dv90 for manual actuations. Each of these parameters met acceptance criteria of 75%-125% of the average for manual actuations. Students' t-tests, while not necessarily appropriate for the small data sets, indicated that the data sets for manual actuations and automated actuations were equivalent for Dv10, but not for Dv50 or Dv90. It was not possible to accurately replicate the droplet size distribution from manual actuations with the MightyRunt. The force required to reliably break the tabs and actuate the devices produced a distribution with smaller droplets than that of the manual actuations. Less aggressive actuation parameters, which should produce larger droplet sizes, were not sufficient for consistent 55 actuation of the devices. Spraytec method development proceeded with these parameters despite not meeting the acceptance criteria specified in the protocol.

Method Development

Method development involved a vignetting study, device placement study and exhaust placement study utilizing the Spraytec. Actuations were performed using the MightyRunt and the previously qualified parameters. For all tests, the device was placed in front of the laser beam path, the plume traveled through the laser path, and the plume was collected in an exhaust manifold placed behind the laser beam.

The vignetting experiments were performed with the device aligned with the front of the instrument (approximately 10 cm from the beam) and varying distances for the device-to-range lens placement. The results are summarized in Table 42.

TABLE 42

Distance to Range Lens (cm)	Dv10 (μm)	Dv50 (μm)	Dv90 (μm)
6	17.2	38.6	65.9
	16.1	36.5	63.0
8	18.6	41.3	71.6
	21.9	44.2	84.6
10	16.7	38.7	80.4
	16.9	39.8	68.0
12	22.1	46.1	89.8
	17.1	37.1	65.5
	15.6	35.7	62.7
14	17.4	42.3	72.5
	16.6	36.5	64.2

A plot of the Dv10, Dv50, Dv90, and plume records values versus placement is set forth in FIG. 11. The data showed no significant trend over the entire range of placements. To minimize the possibility of deposition of droplets on the range lens during testing, the placement of 14 cm (approximately centered between the laser and range lens supports) was chosen and used for all further testing.

To evaluate exhaust placement on the plume during testing, PSD (particle size diameter) data was collected and evaluated at four distances for exhaust-to-laser beam. The data are summarized in Table 43 below.

TABLE 43

Exhaust Height (cm)	Dv10 (μm)	Dv50 (µm)	Dv90 (μm)
7 cn	n Device-to-Laser	Beam Placement	
No exhaust	14.4	28.1	51.7
	13.3	28.6	54.0
3	19.8	37.1	60.9
	18.1	36.3	61.0
4 cn	n Device-to-Laser	Beam Placement	
No exhaust	17.8	35.0	57.9
	18.7	36.3	60.5
1	14.6	29.4	51.0
	12.6	28.0	56.6
3	15.1	30.0	52.0
	13.6	27.6	49.0
5	14.3	32.8	79.8
	13.4	28.7	51.7
7	11.2	25.5	55.7
	15.9	30.0	50.8

A graphical summary of Dv10, Dv50, and Dv90 values versus placement is included in FIG. 12. There were no significant changes in the Dv10, Dv50, Dv90 or plume duration values for plumes collected without exhaust, or with exhaust 1, 3, 5, or 7 cm behind the beam. The exhaust-behind-laser beam placement was chosen to be 3 cm to reduce the chance of deposition outside the fan housing of sprays performed at a device-to-beam distance of 7 cm.

The method was to include two distances for analysis from the tip of the device to the path of the laser for a more complete characterization of the droplet size distribution. 65 Four distances were evaluated, and results are included in Table 44 below.

54

TABLE 44

Distance (cm)	Dv10 (μm)	Dv50 (μm)	Dv90 (μm)
7	16.7	34.5	67.8
	15.8	33.4	60.0
6	17.8	35.0	57.9
	18.7	36.3	60.5
5	19.1	35.5	58.5
	16.9	32.4	55.1
4	11.8	27.8	57.5
	13.0	28.1	50.3
3	14.4	28.1	51.7
	13.3	28.6	54.0

A graphical summary of Dv10, Dv50, and Dv90 values versus placement are included in FIG. 3. There was no observable trend in the data from various heights. While there is no guidance document for sublingual sprays, the FDA Guidance Document "Bioequivalence and Bioavailability Studies for Nasal Aerosols and Nasal Sprays for Local Action", specifies two distances from 3-7 cm. The larger pair of distances (4 cm and 7 cm) from the beam was chosen in order to characterize a more fully developed plume.

Method Validation

An analyst tested six devices at both the 4 cm and 7 cm distances. A second analyst repeated the analyses at both distances with the next actuations after priming from each device on a second day. Validation results are summarized in Tables 45 and 46 below where the Dv10, Dv50, and Dv90 results were compared.

TABLE 45

	Validation PSD Results at 4 cm Device-to-Laser Beam Placement						
	Device	Dv10 (μm)	Dv50 (μm)	Dv90 (μm)	Shot Weight (mg)	% Average	
ı			Analyst	1			
	1	18.9	31.2	52.3	60.4	76	
	2	16.4	30.8	52.6	78.8	100	
	3	17.2	29.8	51.3	65.0	82	
	4	18.3	31.6	54.2	80.1	101	
	5	13.5	28.5	51.7	81.6	103	
	6	15.0	30.1	52.3	77.2	98	
	Average	16.5	30.3	52.4	73.9		
	% RSD	12.3	3.7	1.9	12.0		
			Analyst	2			
	1	14.2	28.8	57.5	76.8		
	2	11.3	26.8	54.9	76.9		
	3	11.8	27.2	52.2	79.9		
	4	15.4	29.2	52.6	80.0		
	5	12.3	29.2	60.2	76.9		
	6	11.8	26.8	52.2	84.5		
	Average	12.8	27.8	54.9	79.2		
	% RSD	12.8	3.7	6.0	3.8		
	% KSD Overall	14.7	29.0	53.7	76.5		
	Average	14.7	29.0	33.7	70.5		
	Overall	17.9	5.8	5.0	9.0		
	% RSD	17.9	5.0	3.0	9.0		
	Analyst 1	129	109	95			
1	Analyst 1 Average	129	109	93			
	as % of						
	Analyst 2						
	t-stat	3.50	4.20	-1.80			
	t-critical	1.81	1.81	1.81			
	Result	Different	Different	Same			
	Result	Population	Population	Population			
		1 opulation	1 opulation	1 opulation			

Validation	PSD Results	at 7 cm Devi	ce-to-Laser Be	am Placei	ment
File	Dv10 (μm)	Dv50 (μm)	Dv90 (μm)	Shot Weight (mg)	% Average
		Analyst 1-7	cm		
a009 a010 a011 a012 a013 a014 Average % RSD	23.3 16.1 16.0 17.6 20.0 19.6 18.8 14.8	37.0 34.0 34.9 34.0 35.5 36.1 35.3 3.4 Analyst 2-7	58.8 57.3 59.7 57.1 58.0 57.2 58.0 1.8	80.1 76.9 79.0 80.7 82.3 80.6 79.9 2.3	104 99 102 104 106 104
		Allalyst 2-7	CIII		
b009 b010 b011 b012 b013 b015 Average % RSD Overall Average Overall % RSD Analyst 1 Average as % of Analyst 2	11.8 17.0 15.3 13.8 13.3 16.6 14.6 13.8 16.7 18.9	28.7 33.9 31.8 31.1 30.7 32.8 31.5 5.8 33.4 7.3	55.0 61.7 55.8 54.6 55.1 56.7 56.5 4.7 57.2 3.6	72.6 79.8 74.8 79.7 80.8 76.3 77.3 4.2 78.6 3.6	
Average t-stat t-critical Result	2.93 1.81 Different Population	4.25 1.83 Different Population	1.32 1.81 Same Population		

Acceptance Criteria

Analyst 1 shot weight results ranged from 76%-103% of analyst 2 average for the 4 cm distance, and 99%-106% for the 7 cm distance, meeting the acceptance criteria of 75%- 40 125%. The average Dv10 for Analyst 1 was 129% of Analyst 2 for the 4 cm distance and 128% for the 7 cm distance. The average Dv50 for Analyst 1 was 109% of Analyst 2 for the 4 cm distance and 112% for the 7 cm distance. The average Dv90 Analyst 1 was 95% of Analyst 2 for the 4 cm distance 45 and 103% for the 7 cm distance. While Dv10 did not meet acceptance criteria of 75%-125%, this is a measurement of the smallest droplets in the plume and a higher variability is not unexpected. Results for Dv50 and Dv90 met acceptance criteria of 75-125%. Students' t-tests, while not necessarily 50 appropriate for the small data sets, indicated that the data sets for Analyst 1 and Analyst 2 were equivalent for Dv90 at both distances, but not for Dv10 or Dv50 at either distance.

Conclusion

A method for the droplet size distribution analysis by laser diffraction for use with fentanyl sublingual spray was developed and subsequently qualified. Acceptance criteria based on statistical analysis with students' t-test were deemed inappropriate for the small data sets. Method validation acceptance criteria for agreement of Analysts 1 and 2 were determined to be too narrow for the high variability associated with measurement of the smallest droplets in the plume (Dv10). While these criteria were not met, the method was deemed 65 acceptable for use. The method is suitable for use within the operating parameters specified herein.

In Example 17, a study was performed to determine the respirable dose less than 9 µm for fentanyl in 1, 2, 4, 6, and 8 mg/mL fentanyl sublingual spray samples with a working concentration of between 0.1 µg/mL and 5 µg/mL fentanyl in solution. The method used was qualified in compliance with GMP requirements. The sample solutions were determined to be stable over a seven-day period in volumetric glassware and amber HPLC vials at refrigerated and ambient conditions.

The HPLC process was consistent with the process described in Example 15 above. The materials and supplies utilized in the study included acetonitrile (HPLC Grade), potassium phosphate monobasic (ACS Grade), phosphoric acid (ACS Grade), deionized water, alcohol (ethanol, absolute), Short Stack Andersen Cascade Impactor set-up consisting of a 5-liter expansion chamber, induction port, stages 0, 1, 2, and after filter, a vacuum source, in-line flow meter (Sierra Top-Track or equivalent), VWR Sterile sampling bags, glass fiber filter, 8.1 cm, external calibrated flow meter (Dry-Cal Flow Meter or equivalent), and a pneumatic actuator (Innova Systems Mighty Runt or equivalent).

Solution preparations were prepared according to the methods described in Example 15 above and may be scaled as required.

The extraction solution was 50:50 (95/5, Ethanol/Acetonitrile:Water). For every liter of prepared solution, 475 mL ethanol, 25 mL acetonitrile and 500 mL of water was combined in a suitable container and mixed well. This solution expires after one month.

The phosphate buffer solution was prepared in a concentration of 50 mM $\rm KH_2PO_4$ with a pH 2.8. For every liter of prepared solution, 6.8 g potassium phosphate mono basic and 1 liter of water is combined in a suitable vessel and mixed well. The pH of the solution was adjusted to pH 2.8 with the drop-wise addition of phosphoric acid. The solution was filtered through 0.45 μm nylon. This solution expires after one month

The impactor set-up will consist of a 5-liter expansion chamber, induction port, stages 0, 1, 2, and filter prepared according to the following procedure.

The filter stage was placed onto the impactor base.

An 8.1 cm glass fiber filter was placed into the after filter stage and secured with a clean rubber o-ring.

A solid plate was placed on top of the filter stage and then stage 2 was placed in position.

A plate with center hole cutout was placed on top of stage 2 and then stage 1 was placed in position.

A plate with center hole cutout was placed on top of stage 1 and then stage 0 was in position.

The cone was placed in position and the impactor was secured with the hold down clamps.

The induction port was affixed to the cone and the 5 L spansion chamber was placed on top of the induction port. Testing Set-up

The testing instrumentation was set up by placing an in-line flow meter between the vacuum source and cascade impactor with appropriate tubing. A leak test was performed on the impactor. A flow through the impactor was started by opening the vacuum source and the flow was adjusted to approximately 28.3 L/min. A hand was placed over the spray actuation port on the expansion chamber. The flow rate as indicated on the in-line meter was expected to fall to zero. If a flow was still registered, the condition of the impactor a-rings was checked, and the test repeated. To set the flow rate, the expansion chamber was removed from the induction port and an

external calibrated flow meter was attached to the induction port and the flow was started. The flow was adjusted to 28.3±1 L/min with the external calibrated meter and the measurement displayed on the in-line flow meter was recorded for using during the testing procedure.

Testing Procedure

Two devices were actuated for the 1 mg/mL product strength for each assay result. One device was actuated for the 10, 4, 6, and 8 mg/mL product strength. With the expansion chamber in place, the vacuum was started and the flow adjusted to the measurement obtained in during the testing set-up.

The pre-actuated weight of the device in grams to a minimum of 4 decimal places was recorded. The device was positioned so that the spray would travel directly toward the wall opposite the actuation port. The sublingual spray device was actuated into the expansion chamber with the automated pneumatic actuator method parameters listed in Table 47.

TABLE 47

Parameter	Setting
Actuation Force (kg)	5.0
Force Rise Time (s)	0.1
Hold Time (s)	1.0
Force Fall Time (s)	1.0
Spray Delays (s)	1
Minimum Travel Distance (mm)	10.0
Maximum Travel Time (s)	4.0
Trigger Signal Delay (s)	0.0
Stage	Yes

The post actuated weight of the device in grams is recorded 35 to a minimum of 4 decimal places.

Extraction Procedure

Extraction was accomplished by breaking down the impactor set-up and extracting each component by the following procedures:

The plates and the filter were each extracted separately in bags with 10.0~mL of extraction solution. The sample was extracted by hand shaking and kneading for at least one minute. 6.0~mL of the extracted sample was transferred to a 10~mL volumetric flask and diluted to volume with phosphate buffer solution and mixed well. This is the sample solution. This solution expires after seven days.

The expansion chamber is inverted and place in a holder. The induction port and cone is inserted into the expansion chamber ground glass joint. Approximately 40 mL of extraction solution is rinsed through the cone and induction port into the expansion chamber. The solution is swirled in the expansion chamber in an effort to extract the entire interior surface. The solution is decanted into a 200 mL volumetric flask. The rinse is repeated two additional times for a total rinse volume of approximately 120 mL. The flask is brought to volume with phosphate buffer solution and mixed well. This is the sample 6 solution. This solution expires after seven days.

Assay

The sample solutions are assayed as per the procedure set 65 forth in Example 15 above. Calculations are performed as follows.

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Response Factor
$$(Rf) = \frac{\text{Fentanyl Peak Area}}{\text{Fentanyl Concentration } (\mu g/\text{mL})}$$

The fentanyl in the plate and filter samples were determined according to the following calculation:

The amount of fentanyl in
$$\mu g = \frac{A_{scamp} * D_f}{WSI R_f OTR}$$

 A_{samp} =Area of fentanyl in sample preparation

D_f=Dilution factor of fentanyl sample solution preparation (10.0 mL×10.0 ml/6.0 mL)

WSI R_fOTR=Working standard I response factor over the run
The fentanyl in the cone, induction port and expansion
chamber is determined according to the following calculation:

Amount of fentanyl in
$$\mu g = \frac{A_{samp} * V_{samp}}{WSI R_f OTR}$$

 $_{25}$ A_{samp} =Area of fentanyl in sample preparation V_{samp} =Volume of fentanyl in sample solution preparation (200 mL)

WSI R_f OTR=Working standard I response factor over the run. The respirable dose for 2, 4, 6, and 8 mg/mL is calculated as follows:

Respirable dose in μg=Sum of the Drug Mass in Particle Size Fraction Less Than 9 μm (μg)

The respirable dose for 1 mg/mL is calculated as follows:

Sum of the drug mass in particle size fraction less than 9
$$\mu$$
g (μ g) =
$$\frac{\text{than 9 } \mu\text{g }(\mu\text{g})}{\text{2 (number of actuations)}}$$

The respirable fraction is calculated as follows:

$$\frac{\text{Percent (\%)}}{\text{Respirable fraction}} = \frac{\frac{\text{Drug mass in particle}}{\text{size fraction less than 9 } \mu\text{m} \ (\mu\text{g})}{\text{Total Drug Mass} \ (\mu\text{g})} \times 100$$

The particle size cutoff diameters for reporting is set forth in Table 48 below.

TABLE 48

	Impactor Component	Particle Size Grouping
55 —	Expansion Chamber, Induction Port, and Cone Plate 0	≧9 µm
50	Plate 1 Plate 2 F	9 μm > X ≧ 5.8 μm <5.8 μm

The results for the fentanyl sample assay results in μg , the respirable dose in the particle size fraction less than 9 μ m (as per Table 48) in μg , and the percent respirable dose less than 9 μ m in percent to one decimal place were reported.

The Certificate of Analysis for the determination of the respirable 1 mg/mL dose set forth in Table 49 below.

TABLE 49

Т	Determination of Respirable Dose, 1 mg/mL Fe				Specification			
Assay of Fentanyl in sublingual spray samples				Described in Example 17			Report Results	
D de	eterminationse in fenta	on of respiration of respirations of respirations of the contraction o	ıble ual			Report R	esults	
CI Run	Sample	Fentanyl (µg/ dose)	Particle Size groupings	Groupings percent	Average Shot weight (mg)	Total Mass <9 μm (μg)	Respirable dose <9 µm (µg)	
1	Globe	76.5694	≧9 μm	96.4	85.4	2.9	3.6	
	Plate 0 Plate 1	0.5479 0.6228	9 μm >X ≧ 5.8 μm	0.8				
	Plate 2 Filter	0.4746 1.8149	<5.8 μm	2.9				
2	Globe Plate 0	78.6941 0.6746	≧9 μm	96.6	84.0	2.8	3.4	
	Plate 1	0.6217	9 μm > X ≧ 5.8 μm	0.8				
	Plate 2 Filter	0.5000 1.6740	<5.8 μm	2.6				
3	Globe Plate 0	78.0529 0.5082	≧9 μm	97.1	85.3	2.3	2.9	
	Plate 1	0.5429	9 μm > X ≧ 5.8 μm	0.7				
	Plate 2 Filter	0.4185 1.3596	<5.8 μm	2.2				
Average percent respirable dose							3.3	

Many other variations of the present invention will be apparent to those skilled in the art and are meant to be within the scope of the claims appended hereto, including but not limited to the particular unit dose or bi-dose devices and the particle size range of fentanyl produced, as well as other numerical parameters described in the examples, and any combination thereof.

What is claimed is:

1. A unit dose of a non-propellant sublingual fentanyl formulation comprising discrete liquid droplets of an effective amount of fentanyl and a pharmaceutically acceptable liquid carrier, wherein the sublingual fentanyl formulation comprises:

from about 0.1% to about 0.8% by weight of fentanyl or a pharmaceutically acceptable salt thereof;

- from about 20% to about 60% by weight of ethanol; and from about 4% to about 6% by weight of propylene glycol;
- wherein after sublingual administration to a human, said sublingual fentanyl formulation provides a mean time to maximum plasma concentration (T_{max}) of fentanyl of from about 5 to about 120 minutes.
- 2. The unit dose of claim 1, wherein said discrete liquid droplets have a size distribution of from about 10 μ m to about 200 μ m.
- 3. The unit dose of claim 1 wherein after sublingual administration to a human, the sublingual fentanyl formulation provides a mean time to maximum plasma concentration (T_{max}) of fentanyl of from about 10 to about 60 minutes.

* * * * *